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The British
Mycological Society

(Recognosce notum, ignotum inspice)

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Edited by

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THE MATLOCK BATH FORAY

31 May-3 June 1935

By J. RAMSBOTTOM

A PARTY of fifteen assembled on Friday, 31 May, at the Matlock Bath Hotel. It was obvious that six weeks without rain had had a very adverse effect on the limestone soil from the point of view of a mycologist.

Cars were taken on the Saturday morning to the Ashbourne end of Dovedale, much of which is now the property of the National Trust. The day was ideal for a stroll by the Dove beloved by Isaac Walton and Charles Cotton, and the eyes of the fishermen were frequently on the stream "full of very good trout and grayling", as Piscator tells Viator. A fair number of rusts were found, some in good quantity, and *Synchytrium Mercurialis* occurred abundantly along the side of the path bordering the Dove.

The Sunday foray was to the woods on both sides of the curiously named Via Gellia,* the upper part of the valley being worked. Hardly had collecting started than the much-needed rain came down in torrents. Agarics were practically absent, and it soon became impossible to continue searching for microfungi in comfort or with profit. On Monday the sky was overcast, but the rain kept off until late in the afternoon. The middle part of Via Gellia (Middleton Wood) was visited, and an intensive search was made for microfungi. All the members present contributed to the subjoined list, particularly Dr Malcolm Wilson and Dr Alex Smith with rusts, and Mr T. Petch and Mr C. G. C. Chesters with Pyrenomycetes; Miss F. L. Stephens worked out some of the microfungi.

* "The road was built by Philip Gell, of Hopton Hall, near Wirksworth, about 1800, to open up the mineral resources of that part of his estates. It was named after the centurion, Phillipus Gellius, who in the early part of the fourth century was in charge of the band of Roman lead-miners on the surrounding hills. His inscribed urn, with the date, was found at Ivett's Low, just above the Via Gellia. Adjacent hills are named Soldier's Knoll and Chariot Clump, and are connected by Chariot Way.

"The name of Phillipus Gellius is also found on the Roman Wall (where he was at one time in charge of a cohort)." (*The Observer*, 19 Jan. 1936.)

"The name Viyella, given to a much advertised mixture of woollen and cotton which is a product of Derbyshire mills, is—*horresco referens*—a corruption of Via Gellia. Its only excuse is that it was perpetrated to satisfy the requirements of the Patent Office." (J. B. Firth.)

List of Species gathered during the Foray

D. = Dovedale; *G.* = Via Gellia; *MB.* = Matlock Bath.

HYMENOMYCETES

- Armillaria mellea* (Vahl) Fr., rhizomorphs only, *G.*
Tricholoma imbricatum Fr., *G.*
Omphalia umbellifera (Linn.) Fr., *G.*
Hypholoma fasciculare (Huds.) Fr., *D.*, *G.*
Claudopus variabilis (Pers.) W. G. Sm., on *Epilobium*, *G.*
Polyporus varius Fr., *D.*, *squamosus* (Huds.) Fr., *G.*, *hispidus* (Bull.) Fr., *G.*,
radiatus (Sow.) Fr., *D.*, *caesius* (Schr.) Fr., *G.*
Fomes pomaceus (Pers.) Big. & Guill. on *Prunus spinosa*, *G.*
Ganoderma applanatum (Pers.) Pat., Cromford.
Polystictus versicolor (Linn.) Fr., *D.*, *G.*
Irpex obliquus (Schr.) Fr., *G.*
Grandinia farinacea (Pers.) Bourd. & Galz., *G.*
Phylacteria terrestris (Ehrh.) Big. & Guill., *G.*
Stereum rugosum (Pers.) Fr., *G.*, *hirsutum* (Willd.) Fr., *D.*, *G.*
Hymenochaete rubiginosa (Dicks.) Lév., *G.*
Corticium Sambuci (Pers.) Fr., *G.*
Peniophora pallidula Bres., *G.*, *byssosidea* (Pers.) v. Hoehn. & Litsch., *G.* *cinerea*
 (Fr.) Cke, *G.*, *quercina* (Pers.) Cke. on *Fraxinus*, *D.*
Cyphella capule (Holmsk.) Fr., *G.*
Pistillaria micans (Pers.) Fr., *G.*
Tremella mesenterica (Retz.) Fr., *G.*
Exidia Thuretiana (Lév.) Fr., *G.*
Dacryomyces deliquescens (Bull.) Duby., *G.*

GASTEROMYCETES

- Lycoperdon pyriforme* (Schaeff.) Pers., *G.*

UREDINALES

- Uromyces Valerianae* (Schum.) Fuck., *D.*, *Alchemillae* (Pers.) Lév., *G.*, *Ficariae*
 (Schum.) Lév., *G.*, *Scillarum* (Grev.) Wint., *G.*, *Poa* Rabenh. on *Poa* and
Ranunculus Ficaria, *G.*
Puccinia Centaureae Mart., *D.*, *G.*, *Carduorum* Jacky on *Carduus crispus*, *G.*,
obtegens (Link.) Tul., *D.*, *G.*, *Lapsanae* (Schultz) Fuck., *G.*, *Taraxaci* Plowr.,
D., *G.*, *Celakovskiana* Bub., *G.*, *Menthae* Pers., Hotel Garden, *Saniculae*
 Grev., *D.*, *G.*, *Pimpinellae* (Str.) Mart., *D.*, *Chaerophylli* Purt., *D.*, *G.*,
pulverulenta Grev., *D.*, *G.*, *Violae* (Schum.) DC., *D.*, *G.*, *Chrysosplenii* Grev.
 on *Chrysosplenium oppositifolium*, *D.*, *fusca* (Relh.) Wint., *G.*, *Lychnidearum*
 Link, *G.*, *Caricis* (Schum.) Rebent. on *Urtica*, *D.*, *Lolii* Niels., *G.*, *glumarum*
 (Sch.) Eriks. & Henn. on *Brachypodium*, *D.*, *sessilis* Schneid. on *Phalaris* and
Allium, *D.*, *Winteriana* Magn. on *Allium ursinum*, *D.*, *Antirrhini* Diet. & Holw.,
 Hotel Garden.
Triphragmium Ulmariae (Schum.) Link, *G.*, *D.*
Phragmidium Sanguisorbae (DC.) Schroet., *D.*, *mucronatum* (Pers.) Schlecht.,
D., *violaceum* (Schultz) Wint., *D.*, *G.*
Melampsora Rostrupii Wagn., *D.*, *G.*
Melampsorella Caryophyllacearum (DC.) Schroet. on *Cerastium arvense*, *G.*
Milesia Scolopendrii (Fuck.), Hotel Garden.
Kuehneola albida (Kuehn.) Magn., *G.*

USTILAGINALES

- Ustilago violacea* (Pers.) Wint., *D.*, *G.*
Entyloma Ranunculi (Bonord.) Schroet., *G.*
Urocystis Anemones (Pers.) Schroet., *G.*

PYRENOAMYCETES

- Sphaerotheca pannosa (Wallr.) Lév., *G.*
 Erysiphe Cichoracearum DC. on *Arctium*, *D.*, Polygoni DC. on *Heracleum*, *D.*
 Uncinula Aceris (DC.) Sacc., *D.*
 Nectria cinnabarina (Tode) Fr., *MB.*, coccinea (Pers.) Fr. on *Sambucus*, *D.*, *G.*,
 on *Ulmus*, *G.*, sinopica Fr. on *Hedera*, *D.*, mammoidea Phill. & Plowr. on
Ulmus, *G.*, Veuillotiana Sacc. & Roum. on *Ilex*, *G.*, episphaeria Fr. on
Diatrype Stigma on *Crataegus*, *D.*
 Chaetomium elatum Kunze, *G.*
 Leptospora ovina (Pers.) Fuck. on *Urtica*, *D.*
 Melanomma pulvis-pyrius (Pers.) Fuck. on *Crataegus*, *D.*, *G.*, on *Ulmus*, *G.*
 Stigmatea Robertiana Fr., *D.*
 Didymosphaeria fenestrans (Duby) Wint. on *Epilobium*, *D.*
 Leptosphaeria acuta (Moug. & Nestl.) Karst. on *Urtica*, *G.*
 Ophiobolus porphyrogonus (Tode) Sacc., *D.*, Bardanae (Fuck.) Rehm on
Heracleum, *D.*
 Quaternaria dissepata (Fr.) Tul., *G.*
 Eutypa Acharii Tul., *G.*
 Diatrypella verruciformis (Ehrh.) Nitschke on *Corylus*, *D.*, nigroannulata (Grev.)
 Nitschke on *Corylus*, *D.*, *G.*
 Diatrype Stigma (Hoffm.) Fr. g on *Crataegus*, *Corylus*, *D.*
 Hypoxylon fuscum (Pers.) Fr., *D.*, multifforme Fr., *D.*
 Xylaria Hypoxylon (Linn.) Grev., *G.*, polymorpha (Pers.) Grev., *G.*
 Scirrha rimosa (Alb. & Schw.) Fuck., *D.*
 Rhopographis filicinus (Fr.) Nitschke.

HYSTERIALES

- Lophodermium Pinastri (Schräd.) Chev., *G.*

DISCOMYCETES

- Ascomyces alnitorquus (Tul.) Sadeb., *G.*
 Calycella citrina (Hedw.) Qué!, *D.*
 Coryne sarcoides (Jacq.) Tul., *D.*
 Orbilia leucostigma Fr., *D.*, xanthostigma Fr., *G.*
 Chlorosplenium aeruginosum (Oeder) de Not. (sterile) on *Fraxinus*, *G.*
 Helotium herbarum (Pers.) Fr., *D.*
 Dasyscypha virginea (Batsch) Fuck., *D.*
 Trichoscypha calycina (Schum.) Boud., *G.*
 Mollisia cinerea (Batsch) Karst., *G.*
 Catinella olivacea (Batsch) Boud., *G.*
 Phacidium multivalve (DC.) Kunze & Schm., *G.*
 Rhytisma acerinum (Pers.) Fr., *G.*

PHYCOMYCETES

- Synchytrium Taraxaci de By. & Woron., *D.*, *G.*, Mercurialis Fuck., *D.*, *G.*
 Entomophthora dipterigena Thaxt., *D.*
 Plasmopara nivea (Unger) Schroet. on *Aegopodium*, *Angelica*, *Myrrhis*, *D.*, *G.*
 Bremia Lactucae Regel on *Centaurea*, *G.*
 Peronospora Alsinearum Casp. on *Cerastium*, *G.*, Ficariae Tul., *G.*, sordida Berk.
 on *Mimulus*, *G.*, parasitica (Pers.) Tul. on *Arabis*, *Sisymbrium*, *D.*

DEUTEROMYCETES

- Actinonema Rosae (Lib.) Fr., *MB.*
 Phoma acuta Fuck., *D.*, siliquastrum Desm., *G.*, samarum Desm., *D.*
 Microphoma Fraxini Delacr., *D.*

Diplodia inquinans West., *G.*

Septoria Bromi Sacc., *D.*, *Fraxini* West., *D.*, *G.*, *Hederæ* Desm., *G.*, *caricola* Desm., *G.*

Cercospora Mercurialis Passer., *D.*, *G.*

Botrytis cinerea Pers., *D.*

Ovularia obliqua (Cke.) Oud., *D.*, *duplex* Sacc., *G.*, *primulana* Karst., *G.*

**Ramularia Colesporii* Sacc., *D.*, *Scolopendrii* Fautr., *MB.*, *Urticæ* Ces., *D.*, *lychnicola* Cke., *G.*, *lactea* (Desm.) Sacc., *D.*, *Spirææ* Peck, *D.*, *G.*, *Tulasnei* Sacc., *G.*, *Gei* (Eliass) Lindr., *D.*, *G.*, *Primulæ* Thüm., *D.*, *montana* Speg., *D.*, *arvensis* Sacc., *G.*, *calcea* (Desm.) Ces., *D.*, *G.*, *Ajugæ* (Niessl) Sacc., *D.*, *G.*, *Stachydis* (Passer) Massal., *D.*, *purpurascens* Wint., *D.*, *Cirsii* Allesch., *G.*, *Lampsanæ* (Desm.) Sacc., *G.*, *Taraxaci* Karst., *D.*, *G.*

Cladosporium herbarum (Pers.) Link, *D.*

Cladotrichum triseptatum B. & Br., *G.*

Ozonium auricomum Link, *D.*

MYXOMYCETES

By T. PETCH

Didymium squamulosum Fr., *D.*

Comatricha nigra Schroet., *D.*, *MB.*

Trichia contorta Rost., *D.*, *varia* Pers., *G.*

Hemitrichia Karstenii List., *G.*

Perichaena corticalis Rost., *G.*, *depressa* Libert, *G.*

Lycogala epidendrum Fr., *G.*

Reticularia Lycoperdon Bull., *G.*

THE TOTNES FORAY

23 to 28 September, 1935

By E. M. WAKEFIELD

THE thirty-ninth Autumn Foray and Annual General Meeting was held at Totnes, South Devon, from 23–28 September 1935, with headquarters at the Seymour Hotel. About thirty members and friends took part.

The first expedition, on Tuesday, 24 September, was arranged under the guidance of Mr W. E. Hiley to Mr Kitson's estate at Hea-tree, on Dartmoor. Unfortunately the day turned out wet and misty, so that little of the moor itself could be seen. The woods were partly pure conifer and partly mixed, and the first wood visited provided a certain amount of shelter, so that vigorous collecting could be carried out. Among many species recorded were *Nolanea icterina*, *Boletus tridentinus*,* *Aleurodiscus amorphus*, and *Empusa apiculata*, the latter reported later by Mr Petch and new to Britain. After lunch a move was made to some higher ground with mixed woods, but conditions were so wet that most people were glad to get back as soon as possible to the cars and make for tea, which was obtained at Manaton. Before leaving members expressed their sincere thanks to Mr Kitson, Junior, who had accompanied them as an additional guide, for the opportunity given to explore these beautiful woods.

Wednesday proved better as regards weather, fine and warm, and was occupied in a most interesting excursion to the estate of Dartington Hall, near Totnes. Here there was considerable variety of ground, including a little pasture and roadside as well as woodland. Almost at once *Puccinia Iridis* was noted on *Iris foetidissima*, which was growing in abundance along a field path. Another noteworthy rust was *Kuehneola albida*, which was common everywhere on bramble. Wood-rotting fungi received some attention: *Ganoderma applanatum* on beech and *Polyporus dryadeus* on oak were among the early finds. Then, later, special attention was drawn to the unusual sight of *Armillaria mellea* fructifications arising about 3 ft. up the trunk of *Pyrus Aucuparia*. Mr Cartwright secured *Coniophorella olivacea*, and Mr Wilkins *Cantharellus amethysteus*. Mr Petch subsequently reported finding *Solenia fasciculata* and *Gonatorrhodiella parasitica* Thaxt., another new British record. At the end of a very pleasant and busy day the party were

* Mr Ramsbottom was shown this species by the late Abbé Bresadola in 1925; it is undoubtedly the same as *Boletus aurantiporus* Howse.

entertained at tea at Dartington Hall, and afterwards heard a most interesting account, given by Dr Slater, Managing Director, of the history, development and aims of this experiment in rural reconstruction. Before leaving they passed a very hearty vote of thanks to the Trustees and staff for their hospitality, and expressed the hope that they might be allowed to repeat the visit at some future time.

The following day, Thursday, good hunting took place in the grounds of Berry Pomeroy Castle, the property of the Duke of Somerset. This proved the richest ground of the week, and many rare species were recorded, among others *Inocybe rhodiola* and *Psalliota xanthoderma* var. *leptotoides*, which many members thought should be regarded as a species, seeing that it is very different in appearance from the usual *P. xanthoderma*, and in fact strongly suggests a *Lepiota*. *Clavaria aurea*, *Xylaria longipes*, *Nectria Ralfsii*, and *N. sinopica* may also be mentioned. *Aleurodiscus aurantius*, which was found in some quantity on a pile of old branches, has not previously been published as British, but was found some years ago near Newton Abbot by Miss Eyre.

Friday morning was occupied in work at headquarters, but in the afternoon a visit was paid to Seale-Hayne Agricultural College, near Newton Abbot, at the kind invitation of the Principal and staff. Mr Beaumont and Dr Gregory took charge of the party, and after a short account of the history of the college various exhibits illustrating the mycological work there were inspected. A walk round the grounds added a few species to the list, but nothing of any importance. After tea and a very hearty vote of thanks to our hosts the party returned to Totnes.

During the week evening meetings were held as usual. At the Annual General Meeting held on Tuesday, 24 September, the officers for 1936 were elected. Mr F. G. Gould was chosen as President for 1936, with Dr Malcolm Wilson and Mr Cartwright as Vice-Presidents.

The other officers were re-elected. New members of Council, elected to replace three members retiring under the rules, were Miss D. M. Cayley, Dr G. Morgan, and Mr W. C. Moore. Mr Chesters was appointed as delegate to the British Association at Blackpool. The Abbé Bourdot was unanimously elected as an Honorary Member.

After some discussion it was decided to accept the cordial invitation of Irish botanists to visit Killarney for the autumn foray in 1936.

On Wednesday evening the President, Dr Malcolm Wilson, delivered his Presidential Address, on "Some Aspects of Forest Pathology". On Thursday evening Mr Carleton Rea was to have given a paper on the genus *Russula*, but through illness was unable to be present. In his place Miss E. L. Stephens gave a talk on the fungi, particularly Gasteromycetes, of South Africa, and did her best to

persuade members of the British Mycological Society to visit South Africa and help them to name their species. Mr Ramsbottom followed with an account of the International Botanical Congress at Amsterdam, to which Miss Wakefield also added some remarks. At the close of the meeting very hearty votes of thanks were passed to the various landowners whose estates had been visited, and a special vote of thanks to Mr Hiley, manager of the Woodlands Department of Dartington Hall, who had arranged the excursions and obtained the necessary permits.

The following list gives the names of species found during the week, with the exception of a few which have remained as puzzles. For assistance in its compilation the secretary is indebted to all members present, but particularly to Mr Petch, Mr Ramsbottom, Mr Pearson, and Mr Chesters.

Complete List of Species gathered during the Foray

H. = Heatree, Dartmoor; *D.* = Dartington Hall; *B.* = Berry Pomeroy; *S.* = Seale Hayne Agricultural College; *T.* = Totnes neighbourhood.

HYMENOMYCETES

- Amanita phalloides* (Vaill.) Fr., *D.*, *porphyria* (A. & S.) Fr., *D.*, *mappa* (Batsch) Fr., *D.*, *pantherina* (DC.) Fr., *B.*, *spissa* Fr., *D.*, *rubescens* (Pers.) Fr., *H.*, *B.*
Amanitopsis vaginata (Bull.) Roze, *D.*, *B.*, *fulva* (Schaeff.) W. G. Sm., *D.*, *strangulata* (Fr.) Roze, *B.*
Lepiota procera (Scop.) Fr., *D.*, *rhacodes* (Vitt.) Fr., *B.*, *S.*, *excoriata* (Schaeff.) Fr., *D.*, *S.*, *acutesquamosa* (Weinm.) Fr., *B.*, *D.*, *felina* (Pers.) Fr., *B.*, *cristata* (A. & S.) Fr., *H.*, *D.*, *B.*, *castanea* Quél., *B.*, *sistrata* Fr., *D.*, *B.*
Armillaria mellea (Vahl) Fr., *H.*, *D.*, *B.*, *mucida* (Schräd.) Fr., *H.*, *D.*, *B.*
Tricholoma albobrunneum (Pers.) Fr., *D.*, *rutilans* (Schaeff.) Fr., *H.*, *D.*, *B.*, *terreum* (Schaeff.) Fr., *B.*, *cartilagineum* Fr. non Bull., *H.*, *cuneifolium* Fr., *H.*, *saponaceum* Fr., *D.*, *nudum* (Bull.) Fr., *B.*, *polioleucum* Fr., *B.*, *sordidum* (Schum.) Fr., *B.*
Clitocybe nebularis (Batsch) Fr., *H.*, *clavipes* (Pers.) Fr., *H.*, *aurantiaca* (Wulf.) Studer, *H.*, *rivulosa* (Pers.) Fr., *S.*, *cerussata* Fr., *D.*, *infundibuliformis* (Schaeff.) Fr., *D.*, *B.*, *vibecina* Fr., *B.*
Laccaria laccata (Scop.) B. & Br., *H.*, *D.*, *B.*, and var. *amethystina* (Vaill.) B. & Br., *H.*, *D.*
Collybia radicata (Relh.) Berk., *H.*, *D.*, *B.*, *platyphylla* (Pers.) Fr., *H.*, *D.*, *B.*, *fusipes* (Bull.) Berk., *D.*, *maculata* (A. & S.) Fr., *H.*, *distorta* Fr., *H.*, *butyracea* (Bull.) Fr., *H.*, *tuberosa* (Bull.) Fr., *H.*
Mycena pelianthina Fr., *H.*, *B.*, *rubromarginata* Fr., *H.*, *D.*, *pura* (Pers.) Fr., *H.*, *B.*, *Adonis* (Bull.) Fr., *B.*, *galericulata* (Scop.) Fr., *B.*, *polygramma* (Bull.) Fr., *B.*, *ammoniaca* Fr., *H.*, *metata* Fr., *H.*, *amicta* Fr., *D.*, *Iris* Berk., *H.*, *vitis* Fr., *D.*, *B.*, *haematopus* (Pers.) Fr., *H.*, *B.*, *sanguinolenta* (A. & S.) Fr., *H.*, *D.*, *B.*, *galopus* (Pers.) Fr., *H.*, *B.*, and var. *alba* Fl. Dan., *H.*, *B.*, *epipterygia* (Scop.) Fr., *B.*
Omphalia fibula (Bull.) Fr., *H.*, *D.*
Hygrophorus eburneus (Bull.) Fr., *B.*, *cossus* (Sow.) Fr., *B.*, *coccineus* (Schaeff.) Fr., *D.*, *B.*, *conicus* (Scop.) Fr., *S.*
Lactarius insulsus Fr., *D.*, *blennius* Fr., *H.*, *chrysorheus* Fr., *H.*, *deliciosus* (Linn.) Fr., *D.*, *pallidus* (Pers.) Fr., *B.*, *quietus* Fr., *H.*, *B.*, *aurantiacus* (Fl. Dan.) Fr., *D.*, *rufus* (Scop.) Fr., *H.*, *fuliginosus* Fr., *D.*, *serifluus* (DC.) Fr., *H.*, *mitissimus* Fr., *H.*, *D.*, *subdulcis* (Pers.) Fr., *H.*, *B.*

- Russula chloroides* (Krombh.) Bres., *H.*, *lepida* Fr., *D.*, *cyanoxantha* (Schaeff.) Fr., *H.*, *B.*, *fellea* Fr., *H.*, *fragilis* (Pers.) Fr., *D.*, *luteotacta* Rea, *D.*, *atropurpurea* (Krombh.) Maire, *D.*, *xerampelina* (Schaeff.) Fr., *D.*, *lutea* (Huds.) Fr., *B.*, and var. *armeniaca* (Cooke) Rea, *D.*, *B.*, *paludosa* Britz., *D.*, *pectinata* (Bull.) Fr., *B.*
Cantharellus cibarius Fr., *D.*, *B.*, *amethysteus* Quél., *D.*, *lutescens* (Pers.) Fr., *D.*
Marasmius peronatus (Bolt.) Fr., *H.*, *oreades* (Bolt.) Fr., *H.*, *S.*, *globularis* Fr., *B.*, *erythropus* (Pers.) Fr., *B.*, *hariolorum* (DC.) Quél., *H.*, *B.*, *dryophilus* (Bull.) Karst., *H.*, *B.*
Androsaceus rotula (Scop.) Pat., *D.*, *androsaceus* (Linn.) Pat., *H.*
Panus torulosus (Pers.) Fr., *D.*, *B.*, *stipticus* (Bull.) Fr., *D.*, *B.*
Lentinus cochleatus (Pers.) Fr., *D.*, *B.*
Volvaria speciosa Fr., *D.*
Pluteus cervinus (Schaeff.) Fr., *D.*, *salicinus* (Pers.) Fr., *H.*, *nanus* (Pers.) Fr., *D.*
Entoloma rhodopolium Fr., *D.*, *B.*, *nidosum* Fr., *D.*
Nolanea proleteria Fr., *H.*, *B.*, *papillata* Bres., *D.*, *cetrata* (Fr.) Schroet., *H.*, *icterina* Fr., *H.*
Leptonia sericella (Fr.) Quél., *H.*
Clitopilus prunulus (Scop.) Fr., *B.*
Claudopus variabilis (Pers.) W. G. Sm., *H.*, *D.*, *B.*
Paxillus involutus (Batsch) Fr., *H.*
Pholiota erebia Fr., *H.*, *B.*, *squarrosa* (Muell.) Fr., *T.*, *D.*, *adiposa* Fr., *B.*, *mutabilis* (Schaeff.) Fr., *H.*
Inocybe tomentosa (Jungh.) Quél., *H.*, *B.*, *geophylla* (Sow.) Fr., *H.*, *B.*, and var. *lilacina* Fr., *D.*, *descissa* Fr., *H.*, *Godeyi* Gill., *H.*, *Cookei* Bres., *H.*, *D.*, *B.*, *rhodiola* Bres., *B.*, *maculata* Boud., *D.*, *obscura* (Pers.) Fr., *D.*
Astrosporina umbrina (Bres.) Rea, *D.*, *petiginosa* (Fr.) Rea, *D.*
Hebeloma glutinosum (Lindgr.) Fr., *H.*, *mesophaeum* Fr., *B.*, *crustuliniforme* (Bull.) Fr., *H.*, *D.*, *longicaudum* (Pers.) Fr., *B.*
Galera tenera (Schaeff.) Fr., *H.*, *hypnorum* (Schränk) Fr., *H.*, *B.*
Naucoria Cucumis (Pers.) Fr., *B.*, *melinoides* Fr., *B.*, *escharoides* Fr., *H.*
Tubaria furfuracea (Pers.) W. G. Sm., *H.*, *D.*
Flammula sapinea Fr., *H.*, *ochrochlora* Fr., *D.*
Cortinarius (*Phlegmacium*) *cyanopus* (Secr.) Fr., *B.*, *infractus* (Pers.) Fr., *B.*, *caerulescens* Fr., *D.* (*Myxadium*), *elator* Fr., *D.*, (*Inoloma*) *alboviolaceus* (Pers.) Fr., *H.*, *B.*, (*Dermocybe*) *cinnamomeus* (Linn.) Fr., *H.*, (*Telamonia*) *torvus* Fr., *B.*, *hinnuleus* (Sow.) Fr., *D.*, *hemitrichus* Fr., *B.*, (*Hydrocybe*) *duracinus* Fr., *D.*, *castaneus* (Bull.) Fr., *B.*, *decipiens* (Pers.) Fr., *D.*, *acutus* (Pers.) Fr., *B.*, *S.*
Crepidotus mollis (Schaeff.) Fr., *T.*, *D.*, *B.*, *calolepis* Fr., *B.*
Bolbitius fragilis (Linn.) Fr., *D.*, *S.*
Psalliotia xanthoderma Genev., var. *lepiotoides* R. Maire, *B.*, *campestris* (Linn.) Fr., *S.*, *sylvicola* (Vitt.) Fr., *D.*, *B.*, *amethystina* Quél., *D.*, *sylvatica* (Schaeff.) Fr., *B.*, *arvensis* (Schaeff.) Fr., *H.*
Stropharia semiglobata (Batsch) Fr., *H.*, *aeruginosa* (Curt.) Fr., *B.*
Hypholoma fasciculare (Huds.) Fr., *H.*, *B.*, *velutinum* (Pers.) Fr., *H.*, *D.*, *Candolleum* Fr., *B.*, *hydrophilum* (Bull.) Fr., *D.*, *B.*
Psathyra fibrillosa (Pers.) Fr., *H.*
Psathyrella gracilis Fr., *B.*, *atomata* Fr., *H.*, *B.*, *disseminata* (Pers.) Fr., *D.*, *B.*
Panaeolus sphinctrinus Fr., *H.*, *S.*, *campanulatus* (Linn.) Fr., *H.*
Coprinus comatus (Fl. Dan.) Fr., *D.*, *atramentarius* (Bull.) Fr., *D.*, *cinereus* (Schaeff.) Cke, *H.*, *B.*, *micaceus* (Bull.) Fr., *H.*, *B.*, *plicatilis* (Curt.) Fr., *S.*
Gomphidius viscidus (Linn.) Fr., *B.*
Boletus elegans (Schum.) Fr., *H.*, *viscidus* (Linn.) Fr., *H.*, *badius* Fr., *H.*, *bovinus* (Linn.) Fr., *H.*, *chrysenteron* (Bull.) Fr., *H.*, *D.*, *B.*, *subtomentosus* (Linn.) Fr., *H.*, *B.*, *versicolor* Rostk., *B.*, *impolitus* Fr., *D.*, *edulis* (Bull.) Fr., *H.*, *B.*, *calopus* Fr., *D.*, *erythropus* Fr. non Pers., *H.*, *purpureus* Fr., *D.*, *tridentinus* Bres., *H.*, *scaber* (Bull.) Fr., *D.*, *castaneus* (Bull.) Quél., *B.*, *porphyrosporus* Fr., *B.*

- Polyporus squamosus* (Huds.) Fr., *D.*, *B.*, *Schweinitzii* Fr., *B.*, *giganteus* (Pers.) Fr. on hawthorn, *H.*, on beech, *D.*, *dryadeus* (Pers.) Fr., *D.*, *hispidus* (Bull.) Fr., *B.*, *radiatus* (Sow.) Fr., *D.*, *adustus* (Willd.) Fr., *H.*, *D.*, *T.*, *fragilis* Fr., *B.*, *caesius* (Schrad.) Fr., *H.*, *tephroleucus* Fr., *B.*
- Fomes Ribis* (Schum.) Fr. on *Euonymus*, *B.*, *ferruginosus* (Schrad.) Mass., *B.*, *ulmarius* (Sow.) Fr., *D.*, *fraxineus* (Bull.) Fr., on *Ulmus*, *T.*, *annosus* Fr., *H.*, *B.*
- Ganoderma applanatum* (Pers.) Pat., *D.*
- Poria hymenocystis* B. & Br., *H.*
- Polystictus versicolor* (Linn.) Fr., *D.*, *B.*, *abietinus* (Dicks.) Fr., *B.*
- Irpex obliquus* (Schrad.) Fr., *D.*, *B.*
- Lenzites betulina* (Linn.) Fr., *B.*
- Trametes gibbosa* (Pers.) Fr., *D.*, *B.*
- Daedalea biennis* (Bull.) Quél., *D.*, *quercina* (Linn.) Fr., *D.*
- Merulius tremellosus* (Schrad.) Fr., *H.*, *corium* (Pers.) Fr., *D.*, *rufus* (Pers.) Fr., *H.*, *D.*
- Phlebia merismoides* Fr., *H.*
- Coniophorella olivacea* (Fr.) Karst., *D.*
- Hydnum repandum* (Linn.) Fr., var *rufescens* (Pers.) Fr., *D.*
- Mycoleptodon ochraceum* (Pers.) Pat., *B.*
- Acia uda* (Fr.) Bourd. & Galz., *D.*
- Tomentella mucidula* (Karst.) v. Hoehn. & Litsch. (= *Hypochnus roseogriseus* Wakef. & Pears.), *H.*
- Stereum spadiceum* Fr., *H.*, *D.*, *rugosum* (Pers.) Fr., *H.*, *hirsutum* (Willd.) Fr., *D.*, *B.*, *purpureum* (Pers.) Fr., *D.*
- Hymenochaete rubiginosa* (Dicks.) Lév., *B.*, *corrugata* (Fr.) Lév. on *Crataegus*, *H.*, *D.*
- Aleurodiscus amorphus* (Pers.) Rabenh., *H.*, *aurantius* (Pers.) Schroet., *B.*
- Corticium botryosum* Bres., *H.*, *confluens* Fr., *H.*, *porosum* B. & C., *H.*
- Vuilleminia comedens* (Nees) R. Maire, *D.*
- Peniophora Molleriana* (Bres.) Sacc. on *Quercus*, *B.*, *quercina* (Pers.) Cke, *D.*, *cinerea* (Fr.) Cke, *B.*
- Solenia fasciculata* Pers., *D.*
- Clavaria cristata* (Holmsk.) Fr., *H.*, *D.*, *cinerea* (Bull.) Fr., *B.*, *rugosa* (Bull.) Fr., *H.*, *aurea* (Schaeff.) Fr., *B.*, *stricta* (Pers.) Fr., *B.*, *vermicularis* Fr., *H.*
- Pistillaria puberula* Berk., *D.*
- Auricularia mesenterica* (Dicks.) Fr., *D.*, *T.*, *auricula-Judae* (Linn.) Schroet., *B.*
- Tremella mesenterica* (Retz.) Fr., *D.*
- Egidia nucleata* (Schw.) Rea, *D.*, *Thuretiana* (Lév.) Fr., *B.*
- Sebacina incrustans* (Pers.) Tul., *B.*
- Calocera viscosa* (Pers.) Fr., *H.*, *B.*, *cornea* (Batsch) Fr., *H.*, *B.*

GASTEROMYCETES

- Cynophallus caninus* (Huds.) Fr., *H.*
- Phallus impudicus* (Linn.) Pers., *H.*
- Lycoperdon saccatum* (Vahl) Fr., *D.*, *B.*, *depressum* Bon., *S.*, *perlatus* Pers., *H.*, *D.*, *pyriforme* (Schaeff.) Pers., *D.*, *B.*
- Bovisteia plumbea* Fr., *S.*
- Cyathus striatus* (Huds.) Pers., *D.*
- Scleroderma aurantium* Pers., *H.*, *D.*, *B.*
- Sphaerobolus stellatus* (Tode) Pers., *D.*

UREDINALES

- Uromyces Rumicis (Schum.) Wint., *B.*, Poae Rabenh., *B.*
 Puccinia Violae (Schum.) DC., *H.*, Lychnidearum Link, *H.*, *D.*, Circaeae Pers.,
D., *B.*, Umbilici Guép. *H.*, *B.*, Angelicae (Schum.) Fuck., *B.*, Conii (Str.)
 Fuck., *B.*, Saniculae Grev., *B.*, obtegens (Link) Tul., *D.*, Cirsii Lasch, *B.*,
 Lampsanae (Schultz) Fuck., *B.*, Primulae (DC.) Duby, *B.*, Veronicae
 Schroet., *D.*, Antirrhini Diet. & Holw., *T.*, Glechomatis DC., *D.*, Menthae
 Pers., *B.*, Iridis (DC.) Wallr., on *Iris foetidissima*, *D.*, *B.*, oblongata (Link)
 Wint., *D.*, sessilis Schneid., *B.*
 Triphragmium Ulmariae (Schum.) Link, *B.*
 Phragmidium Potentillae (Pers.) Wint., *D.*, Sanguisorbae (DC.) Schroet., *D.*,
 Rubi (Pers.) Wint., *D.*, violaceum (Schultz) Wint., *H.*, *D.*, *B.*, Rubi-idaei
 (Pers.) Karst., *H.*
 Kuehneola alba (Kuehn.) Magn., *D.*
 Coleosporium Senecionis (Pers.) Fr., *T.*, Tussilaginis (Pers.) Kleb., *B.*
 Pucciniastrum Abieti-Chamaenerii Kleb., *D.*, Agrimoniae (DC.) Tranzsch., *D.*,
B., Circaeae (Schum.) Schroet., *D.*
 Melampsora Hypericorum (DC.) Schroet., *D.*, Euphorbiae Cast., *T.*
 Milesina Kriegeriana P. Magn. on *Lastraea dilatata*, *H.*, on *Polypodium*, *H.*

USTILAGINALES

- Ustilago violacea (Pers.) Wint., *D.*
 Sphecelotheca Hydropiperidis (Schum.) de Bary, *D.*, *B.*
 Urocystis Anemones (Pers.) Schroet. on *Ranunculus repens*, *B.*

PYRENOMYCETES

- Erysiphe Cichoracearum DC. on *Arctium*, *H.*, *D.*, on *Plantago*, *D.*, Polygoni DC.
 on *Delphinium*, *T.*, on *Heracleum*, *D.*, tortilis (Wallr.) Fr. on *Cornus*, *B.*
 Uncinula Aceris (DC.) Sacc., *H.*
 Phyllactinia corylea (DC.) Karst. on *Fraxinus*, *H.*, *B.*
 Melanospora parasitica Tul. on *Beauveria Bassiana*, *D.*
 Nectria cinnabarina (Tode) Fr., on *Fagus*, *H.*, *B.*, *D.*, coccinea (Pers.) Fr., *D.*, *B.*,
 sinopica Fr. on *Hedera*, *B.*, Ralfsii B. & Br. on *Acer*, *B.*, galligena Bres. on
Quercus, *D.*, mammoidea Phill. & Plowr., *D.*
 Sphaerostilbe aurantiaca Tul. on *Ulmus*, *D.*
 Hypomyces aurantius (Pers.) Fuck. on *Polystictus versicolor*, *D.*, on *Stereum hirsutum*, *B.*
 Apiocrea chrysosperma (Tul.) Syd. on *Boletus*, *D.*
 Claviceps purpurea (Fr.) Tul. on *Dactylis glomerata*, *D.*
 Cordyceps militaris (Linn.) Fr., *D.*
 Trichosphaeria minima (Fuck.) Wint. on *Fraxinus*, *D.*
 Leptospora ovina (Pers.) Fuck., *D.*
 Melanomma fuscidulum Sacc. on *Fagus*, *H.*, pulvis-pyrius (Pers.) Fuck. on *Fagus*,
Corylus and *Tilia*, *H.*
 Stigmathea Robertiani Fr., *H.*
 Diaporthe taleola (Fr.) Sacc. on *Quercus*, *D.*, leiphaemia (Fr.) Sacc. on *Quercus*, *D.*,
 eres Nitschke on *Hedera*, *D.*
 Cryptosphaeria eunomia (Fr.) Fuck. on *Fraxinus*, *D.*
 Eutypella stellulata (Fr.) Sacc. on *Ulmus*, *D.*
 Quaternaria quaternata (Pers.) Schroet. on *Fagus*, *H.*, *D.*, dissepta (Fr.) Tul. on
Ulmus, *D.*
 Diatrypella verruciformis (Ehrh.) Nitschke on *Fagus*, *H.*, quercina (Pers.) Nitschke
 on *Quercus*, *D.*
 Diatrype Stigma (Hoffm.) Fr. on *Fagus*, *H.*, *D.*
 Ustulina vulgaris Tul., *B.*
 Hypoxylon coccineum Bull. on *Fagus*, *D.*, *B.*, fuscum (Pers.) Fr., *B.*, multiforme
 Fr. on *Corylus*, *H.*, rubiginosum (Pers.) Fr. on *Fagus*, *D.*

Nummularia lutea (A. & S.) Nitschke, *D.*
Daldinia concentrica (Bolt.) Ces. & de Not., *B.*
Xylaria Hypoxylon (Linn.) Grev., *H.*, *B.*, *longipes* Nitschke, *B.*
Rhopographus filicinus (Fr.) Nitschke, *H.*

DISCOMYCETES

Peziza aurantia Pers., *D.*
Lachnea hemisphaerica (Wigg.) Gill., *B.*
Ciliaria scutellata (Linn.) Quél., *H.*
Humaria carbonigena Berk., *D.*
Cudoniella acicularis (Bull.) Schroet., *D.*
Calycella citrina (Hedw.) Quél., *B.*
Bulgaria inquinans (Pers.) Fr., *H.*, *D.* (on *Fagus*).
Chlorosplenium aeruginosum (Oeder) de Not., *D.*, *B.*
Dasyscypha calycina (Schum.) Fuck., *H.*
Hyaloscypha hyalina (Pers.) Boud., *D.*
Mollisia cinerea (Batsch) Karst., *D.*
Phacidium multivalve (DC.) Kunze & Schm., *B.*
Pseudopeziza Trifolii (Biv.-Bern.) Fuck., *D.*
Fabraea Ranunculi (Fr.) Karst., *B.*
Rhytisma acerinum (Pers.) Fr., *H.*, *D.*

HYSTERIALES

Lophodermium Pinastri (Schrad.) Chev., *H.*, *Rhododendri* Ces., *H.*

PHYCOMYCETES

Entomophthora americana Thaxt. on fly, *D.*, *sphaerosperma* Fres. on fly, *D.*,
dipterigena Thaxt. on flies, *H.*, *echinospora* Thaxt. on fly, *D.*
Empusa apiculata Thaxt. on flies and leaf-hopper, *H.*
Spinellus fusiger (Link) v. Tiegh., *D.*

DEUTEROMYCETES

Phomopsis oblonga Trav. on *Ulmus*, *H.*, *D.*
Actinonema Rosae (Lib.) Fr., *T.*
Septoria Violae West., *D.*, *Rubi* West., *D.*, *B.*
Monilia aurea Pers., *H.*, *B.*
Microstroma album (Desm.) Sacc. on *Quercus*, *H.*
Ovularia obliqua (Cke) Oud., *H.*
Sepedonium chrysospermum (Bull.) Fr., *H.*, *B.*
Rhinotrichum repens Preuss, *D.*, *B.*
Beauveria Bassiana (Berk.) Vuill. on weevil, *H.*, on bee, *D.*
Trichoderma viride (Pers.) Fr., *D.*
Gliocladium penicillioides Corda on *Dictydium cancellatum*, *H.*
Cephalosporium coccorum Petch on mealy bug, *T.*, *muscarium* Petch on fly, *D.*
Gonatorrhodiella parasitica Thaxt. on *Hypomyces aurantius*, *D.*
Ramularia sambucina Sacc., *H.*, *Primulae* Thuem., *H.*
Polythrincium Trifolii Kunze, *D.*
Cladosporium herbarum (Pers.) Link on flies, *D.*
Cercospora Mercurialis Pass., *D.*, *B.*
Heterosporium gracile Sacc. on *Iris*, *D.*
Isaria farinosa (Holmsk.) Fr. on pupae, *H.*, *B.*
Hymenostilbe arachnophila (Ditm.) Petch on spider, *B.*
Gibellula araneorum (Schw.) Syd. on spider, *B.*

MYCETOZOA

BY MISS G. LISTER

- Ceratiomyxa fruticulosa* Macbr., *H.*, *D.*, *B.*
Badhamia capsulifera Berk., *D.*
Physarum nutans Pers., *H.*, *D.*, *B.*, *viride* Pers., *B.*
Craterium minutum Fr., *B.*, *leucocephalum* Ditm., *B.*
Leocarpus fragilis Rost., *B.*
Didymium difforme Duby, *D.*, *melanospermum* Macbr. var. *minus* Lister, *B.*,
nigripes Fr., *B.*, and var. *xanthopus* Lister, *D.*, *B.*, *squamulosum* Fr., *B.*,
dubium Rost., *B.*
Stemonitis fusca Roth, *D.*, *B.*, *ferruginea* Ehrenb., *H.*
Comatracha nigra Schroet., *H.*, *D.*, *B.*, *laxa* Rost., *H.*, *typhoides* Rost., *D.*
Lamproderma scintillans Morgan, *B.*
Cribraria vulgaris Schrad., *B.*
Dictydium cancellatum Macbr., *H.*, *B.*
Tubifera ferruginosa Gmel., *B.*
Lycogala epidendrum Fr., *H.*, *D.*
Trichia persimilis Karst., *D.*, *varia* Pers., *B.*, *contorta* Rost., *B.*, *decipiens* Macbr.,
H., *D.*, *B.*, *Botrytis* Pers., *H.*
Hemitrichia vesparium Macbr., *D.*
Arcyria cinerea Pers., *D.*, *B.*, *denudata* Wettst., *H.*, *D.*, *B.*, *incarnata* Pers., *H.*, *D.*

LICHENS OF THE TOTNES DISTRICT

BY W. WATSON

When Totnes was selected as the venue of the 1935 Foray visions of Dartmoor Tors and expectations of seeing species of *Gyrophora*, *Umblicaria*, *Alectoria*, *Stereocaulon* and other lichens of the subalpine rocks were conjured up but Pluvius was too liberal in his welcome. The only official excursion on which the Tors could be visited was the Manaton one and the day was so wet that serious lichenological work was not possible. This loss was to some extent compensated for by a visit with Mr Knight to Lydford but the compensation was far from complete, as is evident by the absence from the following list of the above-mentioned plants and also of such common lichens of the Dartmoor uplands as *Lecanora polytropa*, *Lecidea Dicksonii*, *Cladina sylvatica*, *Cladonia uncialis* and *Baeomyces rufus*.

The Berry Pomeroy woods yielded a fair number of corticolous species and on a limestone wall at Dartington many interesting species characteristic of such a habitat were found. A *Verrucaria* which has not yet been described was found and a number of other plants new to Devon. Mr Lamb succeeded in finding *Bacidia prasinoidea*, another addition from the Kerry flora for south-west England.

Some lichens were so frequent and well distributed that it is needless to specify definite localities for them. These are *Evernia prunastri*, *Parmelia perlata*, *P. caperata* and its forms *sorediosa* and *saxicola*, *P. fuliginosa* and its var. *laetevirens*, *P. saxatilis* and its form *furfuracea*, *P. subaurifera*, *P. conspersa*, *Hypogymnia physodes* and its form *labrosa*,

Lecanora chlorona, *L. rugosa*, *L. allophana*, *L. atra*, *Lecidea parasema* and its var. *flavens*, *L. rivulosa*, *L. contigua*, *Pertusaria jaginea*, *P. pertusa*, *P. dealbata*, *P. leioplaca*, *Ramalina farinacea*, *R. fastigiata*, *Placodium callospium* var. *plicatum* (= *P. flavescens*), *Candelariella vitellina*, *Physcia hispida*, *Cladonia pyxidata*, *C. fimbriata* var. *subulata*, *Peltigera canina*, *P. rufescens*, *P. polydactyla*, *Opographa atra*, *O. vulgata*, *Graphis elegans* and *G. scripta*.

In the following list: *B.* = Berry Pomeroy. *D.* = Dartington. *H.* = Holne. *L.* = Lydford. *M.* = Manaton. *T.* = Totnes. An asterisk denotes that the plant has not previously been recorded for Devon.

- Usnea florida* (L.) Web., *B.*, *L.*, *M.*
var. *hirta* (L.) Ach., *B.*, *M.*
U. ceratina Ach., *B.*, *H.*, *L.*, *M.*
*form *incurviscens* Arn., *M.*
U. dasypoga (Ach.) Nyl., *B.*, *D.*
**U. barbata* (L.) Hoff., emend. Mo-
tyka, *B.*, *M.*
U. articulata (L.) Hoff., *D.*
Evernia furfuracea (L.) Mann., *M.*
*form *scobicina* (Ach.) Cromb., *M.*
Parmelia prolixa (Ach.) Carr., *M.*
var. *isidiostyla* Nyl., *M.*
P. revoluta Flk., *D.*, *H.*
P. tiliacea (Hoff.) Ach., *H.*
P. sulcata Tayl., *D.*
P. dubia (Wulf.) Schaer., *D.* On
rocks, Dartmouth and Strete
P. omphalodes (L.) Ach., *L.*, *M.*,
Dartmeet
Hypogymnia physodes (L.) Nyl.
var. *tubulosa* (Schaer.) Wats., *M.*
Platysma glaucum (L.) Nyl., *B.*, *D.*,
M.
var. *fallax* (Web.) Nyl., *D.*, *M.*
Lecanora albescens (Hoff.) Flk. (= *L.*
galactina Ach.), *L.*, *T.*
L. dispersa (Pers.) Roehl., *T.*
L. Hageni Ach., *M.*, *T.*
L. subfusca (L.) Ach., *B.*, *M.*
L. campestris (Schaer.) Hue, *M.*, *T.*
L. intumescens (Reb.) Krb., *D.*, *H.*, *L.*
L. pallida (Schreb.) Schaer., *H.*
L. carpineae (L.) Wain., *B.*, *D.*
L. expallens (Pers.) Ach., *D.*, *M.*
L. varia (Ehrh.) Ach., *M.*, *T.*
L. symmicta Ach., *H.*, *L.*
var. *sepincola* (Ach.) Nyl., *L.*
L. symmictera Nyl., *D.*, *H.*, *L.*
Ochrolechia parella (L.) Arn., *D.*, *L.*,
M.
*form *porinoides* (Ehrh.) Zahl., *D.*
Aspicilia calcarea (L.) Krb., *B.*, *D.*
A. lacustris (With.) Th. Fr., *H.*
Biatora lucida (Ach.) Fr., *L.*
B. coarctata (Sm.) Th. Fr., *L.*, *M.*
Lecidea macrocarpa (DC.) Steud., *M.*
L. sorediza Nyl., *M.*
**L. cinereoatra* Ach., *M.*
**Pertusaria pertusa* (L.) Tuck.
var. *leiotera* (Nyl.) Zahl., *M.*
P. dealbata (Ach.) Cromb.
form *corallina* (L.) Cromb., *M.*
P. multipuncta (Turn.) Nyl., *D.*
P. globulifera (Turn.) Nyl., *D.*
P. Wulfenii DC., *B.*, *D.*
Biatorella simplex (Dav.) B. & R.,
Strete
Ramalina calicaris (L.) Fr., *B.*, *D.*
R. fraxinea (L.) Ach., *D.*
Solenopsora holophaea (Mont.) samp.,
Dartmouth
Biatorina erysiboides (Nyl.) Th. Fr.,
D., *L.*
B. lenticularis (Ach.) Krb., *D.*
B. Lightfootii (Sm.) Mudd, *B.*, *H.*
Toninia aromatica (Sm.) Mass., Dart-
mouth
Bilimbia sabuletorum (Flk.) Arn., *D.*,
Kingsbridge
Bacidia luteola (Schrader.) Mudd, *B.*
B. phacodes Krb., *L.*
B. effusa (Sm.) Arn., *L.*
**B. prasinoides* (Nyl.) Oliv., *D.*
Leciographa parasitica (Flk.) Mass.,
D.
Xanthoria parietina (L.) Th. Fr., *T.*
var. *ectanea* (Ach.) Oliv., Dart-
mouth
Placodium murorum (Hoff.) DC., *T.*
P. xantholytium Nyl., *D.*
Callospisma citrinum (Hoff.) Mass., *T.*
C. ochraceum (Schaer.) Mass., *D.*
C. aurantiacum (Lightf.) Mass. var.
flavovirescens (Wulf.) Krb., *D.*
C. caesiorufum (Ach.) Wats., Dart-
mouth
C. ferrugineum (Huds.) Mudd, var.
festivum (Ach.) Mudd, *L.*

- C. rupestre* (Scop.) Wats. var. *calvum* (Dicks.) Wats., *L.*
Pseudophyscia fusca (Huds.) Wats., Slapton, Dartmouth
Physcia aipolia (Ach.) Nyl., *T.*
P. virella (Ach.) Lynge, *T.*
Diploicia canescens (Dicks.) Mass., *D.*
Buellia myriocarpa (DC.) Mudd, *D.*
B. stellulata (Tayl.) Mudd, Strete
 **B. verruculosa* (Borr.) Mudd, Strete
B. disciformis (Fr.) Mudd, *H.*
 **B. confervoides* Kremp., Strete
Rhizocarpon geographicum (L.) DC., *M.*, Dartmeet
 **R. viridiatrum* (Flk.) Krb., *L.*, *M.*
R. petraeum (Wulf.) Mass., *L.*, *M.*
Thelotrema lepadinum Ach., *B.*
Phlyctis agelaea (Ach.) Krb., *B.*, *D.*
Diploschistes scruposus (L.) Norm., Dartmeet
Cladonia ochrochlora Flk., *M.*
C. cervicornis (Ach.) Schaer. var. *subcervicornis* Wain. *M.*
C. furcata (Huds.) Schrad., *M.*
C. squamosa Hoff., *L.*
C. rangiformis Hoff., *M.*
C. flabelliformis (Flk.) Wain., *M.*
C. macilentata (Ehrh.) Hoff., *B.*
C. coccifera (L.) Willd., *D.*, *M.*
Coenogonium ebeneum (Dillw.) A. L. Sm., *M.*
Crocynia lanuginosa (Ach.) Hue, *T.*
Lobaria pulmonaria (L.) Hoff., *B.*
L. laetevirens (Light.) Zahl., *B.*
Stictina fuliginosa (Dicks.) Nyl., *L.*
S. limbata (Sm.) Nyl., *L.*
S. Dufourii (Del.) Nyl., *L.*
Peltigera rufescens (Weis.) Humb. var. *praetextata* (Flk.) Nyl., *M.*
 *var. *lepidophora* (Nyl.) Wain., *D.*
P. polydactyla (Neck.) Hoffm. var. *hymenina* (Ach.) Nyl., *B.*, *M.*
P. horizontalis (L.) Hoff., *L.*, *M.*
P. scutata (Dicks.) Krb., *L.*
Pannaria rubiginosa (Thunb.) Del. var. *conoplea* (Pers.) Krb., *L.*
Placynthium nigrum (Huds.) Gray, *D.*
Parmeliella corallinoides (Hoff.) Zahl., *L.*
Leptogium sinuatum (Huds.) Mass., *D.*
 var. *scotinum* (Ach.) Krb., Dartmouth
L. lacerum (Lilj.) Gray, *B.*, *D.*, *L.*
L. ruginosum (Duf.) Nyl., Dartmouth
L. tremelloides (L.) Gray, Dartmouth
L. microscopicum Nyl., *L.*
L. plicatile (Ach.) Th. Fr., *L.*
Collema pulposum (Bern.) Ach., *D.*, *L.*
C. granosum (Scop.) Schaer., *D.*
C. granuliferum Nyl., *D.*
C. cheileum Ach., *D.*, *L.*
 *form *graniforme* Harm., *D.*
Ephebeia hispidula (Ach.) Nyl., *H* (c. fr.).
Arthonia gregaria (Weig.) Krb., *D.*
 *var. *astroidea* (Leight.) Mudd, *L.*
 *var. *cuspidans* (Nyl.) Wats., *L.*
A. lurida Ach., *B.*
 **A. aspersella* Leight., *B.*, *H.*
A. radiata (Pers.) Ach., *B.*
 var. *Swartziana* (Ach.) Sydow, *B.*, *L.*
 (4*) form *parallela* (Mudd) Leight., Okehampton
Opegrapha atra Pers. var. *denigrata* (Ach.) Schaer., *D.*, Okehampton
O. betulina Sm., *B.*
O. calcarea Turn., *M.*
O. saxicola Ach., *D.*
 var. *Decandollei* Stiz., *D.*
O. varia Pers., *D.*
Enterographa crassa (Del.) Fée, *B.*, *D.*
E. Hutchinsiae (Leight.) Krb., *B.*
Graphis scripta (L.) Ach.
 var. *serpentina* (Ach.) Nyl., *D.*
Phaeographis dendritica (Ach.) Mull., *B.*, *D.*
Graphina anguina (Mont.) Mull., *B.*
Calicium hyperellum Ach., *D.*
 **Cyphelium stigonellum* (Ach.) Zahl., *M.*
Coniocybe furfuracea (L.) Ach., *L.*, *M.*
Stenocybe byssacea (Fr.) Nyl., *B.*
Sphaerophorus globosus (Huds.) Wain., *M.*
 form *congestus* (Lamy) A. L. Sm., *M.*
S. fragilis (L.) Pers., *M.*
Dermatocarpon miniatum (L.) Th. Fr., *D.*
Verrucaria maura Wahl., Slapton, Dartmouth
V. viridula (Schrad.) Ach., *D.*
V. maculiformis Kremp., *B.*
V. papillosa Ach., *L.*
V. glaucina Ach., *D.*
V. rupestris Schrad., *D.*
V. nigrescens Pers., *D.*, *T.*
V. muralis Ach., *D.*
V. sphinctrina (Duf.) Nyl., *D.*
 **V. calciseda* DC. emend Stein, *D.*
 *form *calcivora* Mass., *D.*
Acrocordia epipolaea (Borr.) A. L. Sm., *D.*

Arthopyrenia punctiformis (Pers.) Arn., B., H.	*P. affinis (Mass.) Zahl., L.
A. fallax (Nyl.) Arn., H.	*P. leptalea (D. & M.) A. L. Sm., H.
A. submicans (Nyl.) A. L. Sm., B., M.	Pyrenula nitida (Weig.) Ach., B., D.
Porina carpinea (Pers.) Zahl., H., L.	form geographica (Cromb.) Wats., B.
P. chlorotica (Ach.) Wain., D., M.	var. nitidella (Flk.) Mudd, B., D.

Owing to the restricted conception of some of the species of *Usnea* by Dr Motyka of Poland, it seems necessary to mention that the Dartington *U. dasypoga* scarcely agrees with the species as restricted by him, though it agrees with *U. dasypoga* of most authors. On the other hand, *U. florida* var. *hirta*, a name which has been very loosely applied in this country, agrees with both Stirton's and Motyka's conceptions of *hirta*.

NEW GOLD COAST FUNGI. I.

By H. A. DADE

(With 4 Text-figures)

THIELAVIA SETOSA N.SP. (Gold Coast No. 555)

THIS fungus was found clambering over the aerial growths of mixed cultures of moulds growing on several samples of cacao "beans" which had been incubated in a damp atmosphere in Petri dishes. Here it was rendered conspicuous by its very profuse formation of characteristic ascocarps.

DESCRIPTION

When mouldy cacao beans are incubated in a humid atmosphere, a dense growth of Aspergilli forms on their surface, and above these there is commonly a luxuriant tangle of the aerial mycelium and sporophores of *Absidia* spp. In these conditions *Thielavia setosa* grows most vigorously and produces large and fully developed ascocarps. Its mycelium consists of fine hyphae which are strung, like spiders' silk, over and about the *Absidia* hyphae and among the heads of the Aspergilli. On these filaments the ascocarps are borne.

Where less aerial mould growth occurs, as in culture on agar, ascocarps appear on the surface of the medium; and when the competing fungi are predominant, they may be immersed in the medium.

The ascocarps are globose, brilliantly white and shining when young, later becoming brown, and black when mature, due to the dark brown or olivaceous colour of the cortical cells.

The most characteristic morphological feature of the ascocarp is the bristling array of appendages which project radially from all sides, like the spines of a sea urchin or the spicules of *Radiolaria*. These are fully developed in vigorous aerial growths, but may be few and poorly developed in other situations; in immersed ascocarps they are represented only by dark radiating hyphae of unspecialized form. When fully developed, the appendages are simple, rigid and acicular, with expanded and somewhat bulbous bases which fit into the mosaic of cortical cells. They pass through the same colour changes as the ascocarp itself.

In the development of the ascocarp, initials are formed from a short simple spiral of two or three turns, borne on a single hypha. The knot of investing hyphae which soon surrounds and conceals the spiral seems to be derived from one or two small laterals which spring from the stalk of the fertile hypha.

The asci are pear-shaped when young, becoming globose when full grown, with very thin transparent walls. Each contains eight ascospores.

The ascospores are hyaline at first and dark fuliginous or nearly black when mature. At each pole of the spore there is a pore in the exospore, through which the endospore protrudes slightly as a transparent papilla. The papillae cause the broadly fusiform spores to resemble lemons in form.

Germination is generally bipolar, though it may sometimes occur at one pore only. The germ-tubes are expanded where they emerge from the spore, but are not typically lop-sided like those of *Thielavia basicola* described by McCormick (1).

Aleuriospores, which are generally produced, are ovoid and transparent, either sessile on the sides of distributive hyphae, or borne terminally on short laterals. Occasionally they are terminal on long hyphae, the lower cells of which bear sessile spores. Their cell walls are somewhat thickened and refractive, and they may function as chlamydospores. Germination has not been observed.

Repeated search has been made for other spore forms, but none has been found.

CULTURAL CHARACTERISTICS

Attempts to isolate the fungus in pure culture were at first unsuccessful, the separation plates becoming covered by the contaminants before the desired growth appeared. Eventually clean cultivations were obtained by picking out ascocarps which had formed on the reverse side of filter paper used to keep the Petri dishes moist. These pure cultures showed very slow growth and produced only a few ascocarps, none of which contained spores.

One of these pure cultures became contaminated by a species of *Scopulariopsis*, and vigorous growth and an abundant crop of ascocarps ensued. This accident, indicating the Heald-Pool reaction, provided an early clue to successful cultivation, and several "commensals" were tried, *Aspergillus flavus* being the most satisfactory. It was then found that equally good results could be obtained by planting the fungus on slants of potato-dextrose agar previously occupied by young (three to six days old) cultures of *A. flavus* and resterilized by immersing the tubes in boiling water for about half an hour.

This recalled the work on *Thielavia basicola* by McCormick (1), who found that this species could be induced to fruit on media dosed with sterile filtrates containing fungic extracts. The same method succeeds with the present fungus, but the simpler method described above is equally effective, though McCormick found it useless for *T. basicola*. The age of the preliminary cultivation of *Aspergillus flavus* is important. It must be allowed to grow until sufficient of the substances

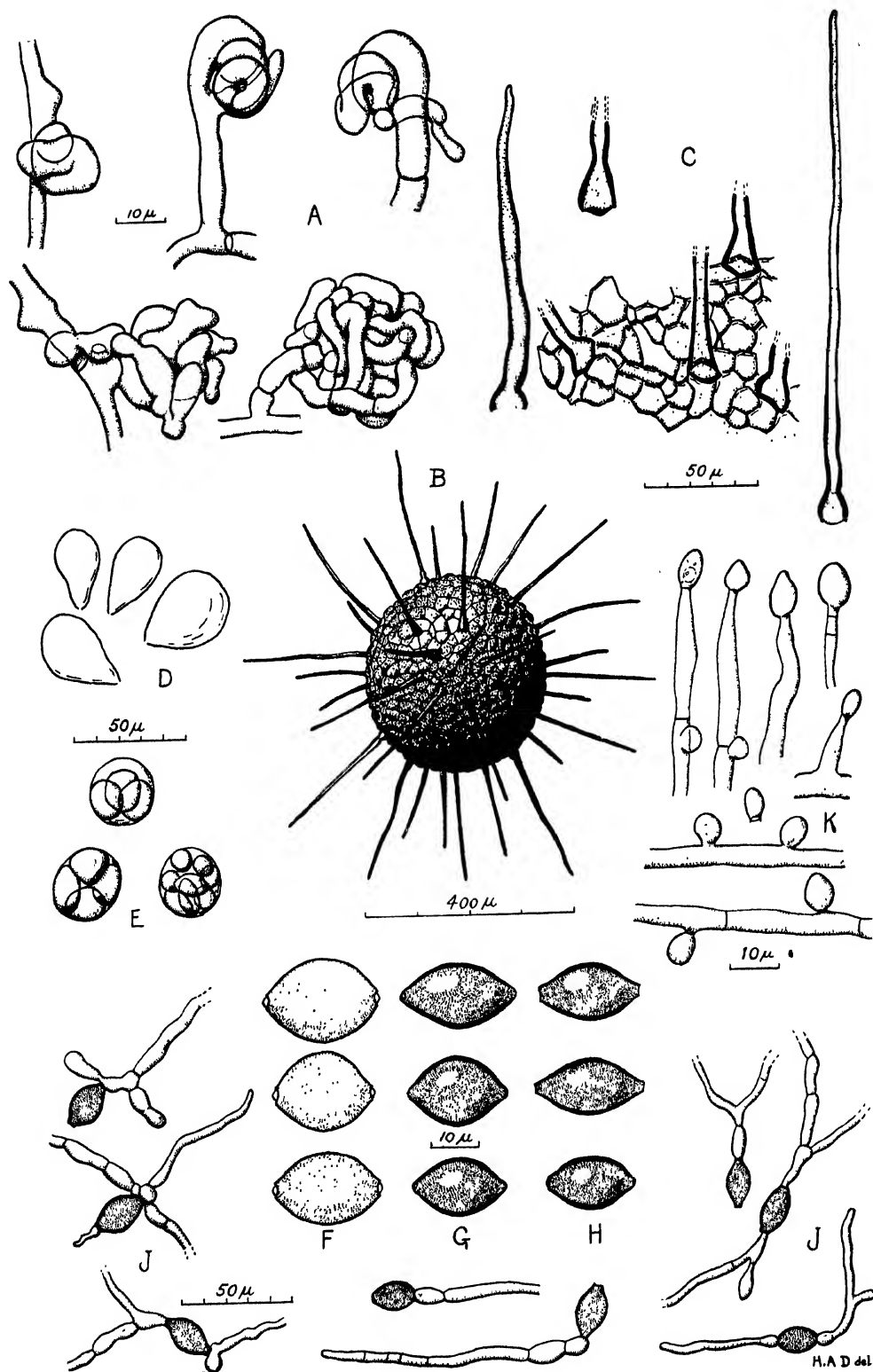


Fig. 1.

necessary for growth of *Thielavia setosa* has accumulated. Very young growths do not give good results. Presumably the extracts of *Aspergillus flavus* required by *Thielavia setosa* are not entirely destroyed by the short heating necessary to kill the former fungus.

In pure culture on virgin medium there is a slow growth of mycelium which forms a fine white or creamy felt. In the more vigorous of these vegetations a few fruit bodies may be formed, hidden within a thick, fluffy felted mass of hyphae, but they are not well developed and contain no spores, and attempts at subculture usually fail. Mass transfers from growths on sterilized *Aspergillus flavus* slants are somewhat more successful, probably because a sufficient quantity of *flavus* extract is carried over with the transfer, but again subcultures do not grow. These observations were made in the Gold Coast at laboratory temperatures ranging from 25 to 30° C. Cultures were sent to Mr R. H. Bunting in England, and in some he found that a good crop of ascocarps, with mature ascospores, was produced in the incubator at a temperature of 24° C. in about three weeks. This may indicate that the effect of temperature is similar to that in *Melanospora Zamiae* Corda, which in the Gold Coast will grow only with a commensal, or in the presence of extracts, like *Thielavia setosa*, while pure cultures sent to England grow freely to maturity with no stimulation by other fungi.

In mixed culture with living growths of the several other fungi which have been tried, *Thielavia setosa* always produces fertile ascocarps, but its reaction with its commensals varies. With *Aspergillus flavus* (typical strain from cacao) the two fungi, planted together at the centre of the plate, advance together, *Thielavia setosa* producing aerial hyphae, which clamber over the heads of the *Aspergillus*, and concentric rings of ascocarps. The fruiting of *A. flavus* is considerably repressed, but its vegetative growth keeps pace with that of its companion. The same proportionate growth occurs with the *Scopulariopsis* sp. previously mentioned. *Aspergillus Chevalieri* and *A. Tamaris* (cacao strains) are, however, completely suppressed by *Thielavia setosa*, and both fungi then cease to extend their centrifugal growth. On the other hand *Aspergillus niger* (cacao strain) can successfully compete with *Thielavia setosa*, the latter producing ascocarps only during the early growth of the dual planting, and sending out a few radial aerial hyphal threads for a short distance, after which it ceases to advance.

It is clear that while the substances produced by all these com-

Legend to Fig. 1.

Fig. 1. *Thielavia setosa* Dade. A, ascocarp initials, $\times 716$ d.; B, fully developed mature ascocarp, $\times 78$ d.; C, setae, and bases showing insertion on ascocarp wall, $\times 330$ d.; D, young asci, $\times 330$ d.; E, asci with developing ascospores, $\times 330$ d.; F, maturing ascospores, from cacao; G, mature ascospores, from cacao; H, mature ascospores, from potato-dextrose agar, $\times 716$ d.; J, germinating ascospores, $\times 330$ d.; K, aleuriospores, $\times 716$ d.

mensals are favourable to *T. setosa*, there is competition, with varying results, for the nutrition provided by the substratum.

Similar reactions occur when the two fungi are planted at some distance from one another on the same plate. Normal pure culture growth occurs until the circumferences of the two vegetations meet. With a favourable commensal the hitherto feeble, sterile *T. setosa* advances rapidly in a fan centred on the point of contact, into the area occupied by the other fungus, and forms a profusion of perithecia. The companion fungus may continue to grow into the *T. setosa* area, but no ascocarps are formed on the sterile mycelium of the latter, where growth has ceased, and which is therefore no longer susceptible to stimulation.

Ascospores germinate quite well in hanging drops of normal potato dextrose and other fluids, but the percentage of germination is distinctly greater in a medium to which fungic extract has been added.

TAXONOMY

This fungus seems to be best included in the genus *Thielavia*, from the other species of which it differs in the presence of well-developed appendages on the ascocarps and in the very large ascospores. The latter are, however, typical of the genus in form. Only one other species—*T. Sepedonium* Emmons (2)—has aleuriospores. In its dependence for development, at least at high temperatures, on the presence of extracts of other fungi, *T. setosa* strongly resembles *T. basicola*.

It is of interest to note that similar nutritional phenomena occur in the genus *Melanospora*. Moreover, the ascospores and terminal chlamydospores of *M. Zamiae* (3) strongly resemble the ascospores and aleuriospores of *Thielavia setosa*. There is also a remarkable resemblance between the young ascocarps of *Melanospora Zamiae* and *Thielavia setosa*. These two genera are undoubtedly closely related.

DIAGNOSIS

Thielavia setosa n.sp.

Ascomata typice aeria vel superficialia nonnunquam immersa; globosa, sine ostiolo, nigra, membranacea, 150–300 μ , plerumque 250–300 μ diam., cum appendicibus multis radiantibus. Appendices rigidae, aciculares, 150–300 μ longae, 8–19 μ crassae, basi dilatae 12–16 μ crassae.

Asci globosi, subtiles, 8 sporas continent.

Ascosporae fuligineae vel subnigrae, citrifformes, 20–25 μ longae, 13–15 μ latae, papillis polaribus hyalinis.

Aleuriosporae ellipticae vel ovatae, hyalinae, ad 8 μ longae, 7 μ latae, sessiles vel in hyphis brevibus lateralibus terminales.

Aliae sporae ignotae.

Saprophila in seminibus *Theobromae Cacao* L. in Gold Coast, Africae Occidentalis.

SYNCEPHALIS NANA N.SP. (Gold Coast No. 570)

This fungus was found as a parasite on a strain of *Absidia Regneri* (Lucet & Cost.) Lendner, which is a mould of inadequately dried cacao beans in the Gold Coast. The samples of cacao beans in which the parasite appeared were from Kpeve, in the British Mandated Territory of Togoland.

In the course of work on cacao moulds, samples of the split beans were incubated on moist filter paper in Petri dishes, and a luxuriant aerial growth of the *Absidia* was obtained. Examination with a binocular magnifier disclosed the presence, among the *Absidia* mycelium, of a small organism with minute yellow heads. This was studied in the original growth and in subsequent cultures, free from other fungi, on artificial media.

The aerial mycelium of the host becomes invested with a fine web of delicate hyphae which form a network, with swollen anastomoses, typical of *Syncephalis*. Fruiting in the *Absidia* is inhibited to some extent, and the stolons and sporangiophores of the host become packed with fine intracellular hyphae.

The fertile hyphae of the parasite form rather short sporangiophores, thick at the base and tapering to the insertion of the inflated apex. A well-developed foot with rather short thick rhizoids firmly clasps the host hypha. The head, or sporangium, is pear-shaped, somewhat flattened on top, and tapering below into the stalk. On the rather flattened apex the sporangium bears between thirty and forty "partial sporangia" (Gäumann's term (4)). These are carried on short sterigmata, which appear as warty protuberances on heads from which the spores have fallen.

The partial sporangia are narrowly ampulliform at first—resembling the phialides of *Aspergillus*—and subsequently take on a form determined by the contained spores. Each contains two spores, shaped like short cucumbers, and set together in a line which follows the curvature of the spores. The concave side of the pair faces inwards, and the complete set of partial sporangia therefore assumes a barrel shape. The colour of the mature spores is that of clear honey. The pairs of spores separate when they are shed. The cluster of partial sporangia frequently falls from the head in a mass, with some tendency to remain adhering in a barrel-shaped bundle for a time.

Zygospores have not been found.

The fungus is easy to maintain in culture with its host, and appears after the *Absidia* has made a good growth. It is a highly specialized parasite, for it does not destroy nor seriously depress its host. Prepara-

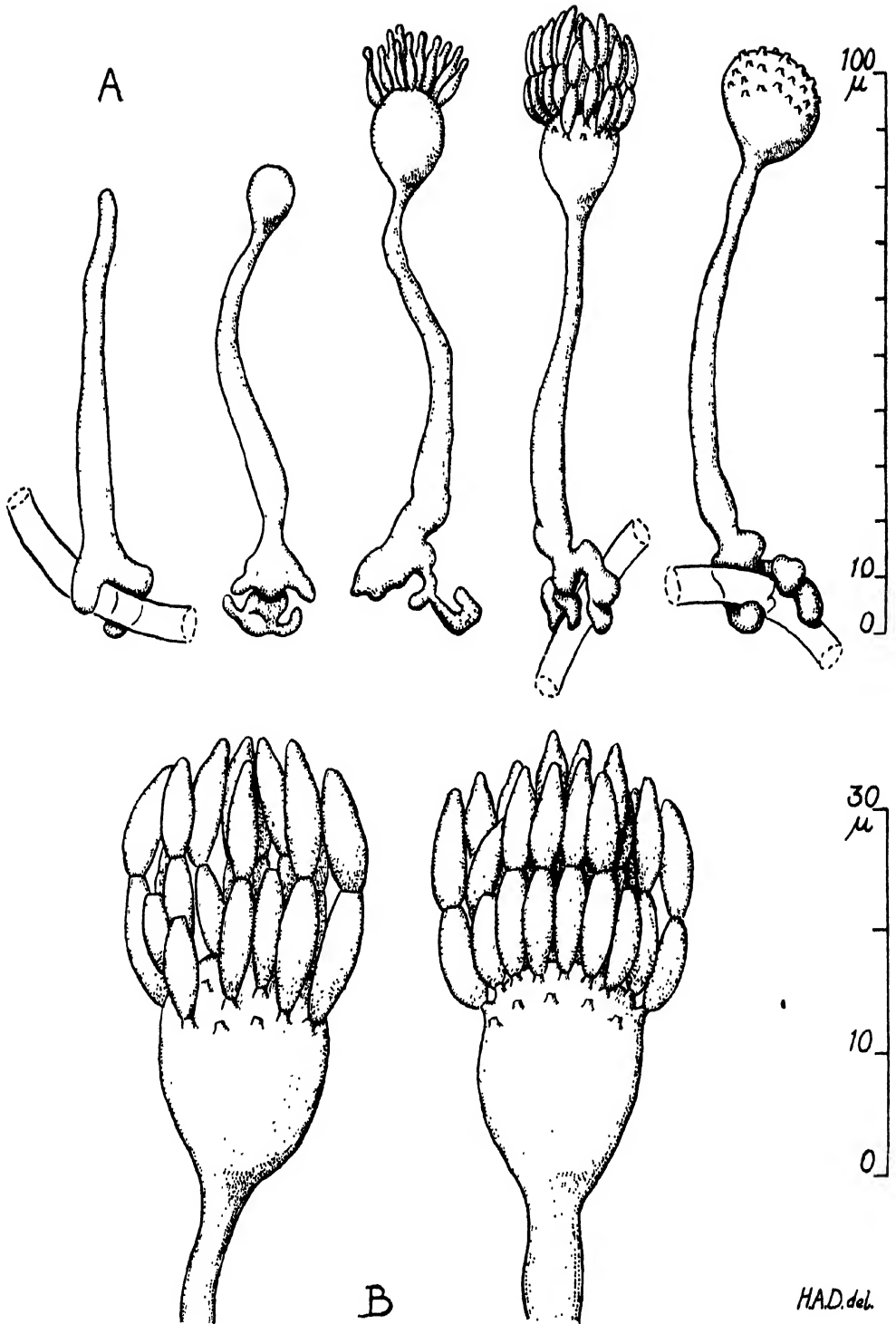


Fig. 2. *Syncephalis nana* Dade. A, complete developing and mature sporangiophores, $\times 660$ d.; B, inflated apices of sporangiophores, bearing clusters of "partial sporangia" and spores, $\times 1432$ d.

tions in Amann's medium, with which cotton blue is incorporated, successfully demonstrate the parasite, for the mycelium, especially the fertile hyphae, takes up the cotton blue more readily than does the *Absidia*.

TAXONOMY

There is no species resembling this fungus among those described by Fischer⁽⁵⁾. The nearest in Fischer's list is *Syncephalis sphaerica*, with undivided "basidialconidien" and rod-shaped spores, but these spores are mostly in five-jointed straight chains. *S. fusiger*, with chains of two spores each, has compound "basidialconidien".

Thaxter⁽⁶⁾ describes a species, *S. tenuis*, with "fertile hyphae septate at base, very elongate, tapering to a slender extremity which expands abruptly to form the fertile head, the latter somewhat flattened and bearing from six to many sporangial filaments arising from all parts of the upper surface or arranged in a more or less definite circle, each producing two spores. Spores subcylindrical to asymmetrically oval, truncate or bluntly rounded, cylindrical forms $20-25 \times 7\mu$, oval forms $25-27 \times 7\mu$ at base to $4-5\mu$ at apex. Sporiferous head without spores $10-20\mu$ diam."

This is much closer to the present fungus, but the latter is smaller and has by no means a "very elongate" stalk. These differences are considered to be sufficient to warrant the recognition of a new species, for which the name *Syncephalis nana* is proposed.

DIAGNOSIS

Syncephalis nana n.sp.

Mycelium cum nodis tumidis reticulatum, typicum generis.

Hyphae fertiles cum capitibus inflatis pedibusque bene formosis, prope basin 8μ diam., apice $2-3\mu$ diam., hyalinae, in maturitate flavidae. Pedes simpliciter ramosi, qui hyphas hospitis complexi sunt. Vesiculae piriformes, $15-20\mu$ crass., in apicibus planulis $30-40$ sporangia ferentes.

Sporangia $16-22\mu$ longa, subcylindrica curvulata, in sterigmatibus brevibus, 2 sporas continent.

Sporae a sporangiis secedentes et saepe in fasciculum doliiformem cohaerentes, fusiformes curvulatae vel cucumeriformes, melleae, $10-12\mu$ long., 4μ crass.

Zygosporae ignotae.

Parasitica in *Absidia Regneri* (Lucet & Cost.) Lendner in Gold Coast, Africae Occidentalis.

MUCOR INAEQUISPORUS N.SP. (Gold Coast No. 637)

At Aburi, in the Gold Coast, ripe fallen fruits of the hog plum, *Spondias monbin* L., soon become covered with a velvety orange-yellow mould. This was found to be a *Mucor* with the following characteristics.

The sporangiophores are typically of *Monomucor* form—i.e. simple unbranched aerial hyphae with terminal sporangia. Rarely, a small lateral branch and sporangium are produced. On fruit of hog plum the fertile hyphae are from 9.5 to 0.75 cm. in length; on potato-dex-

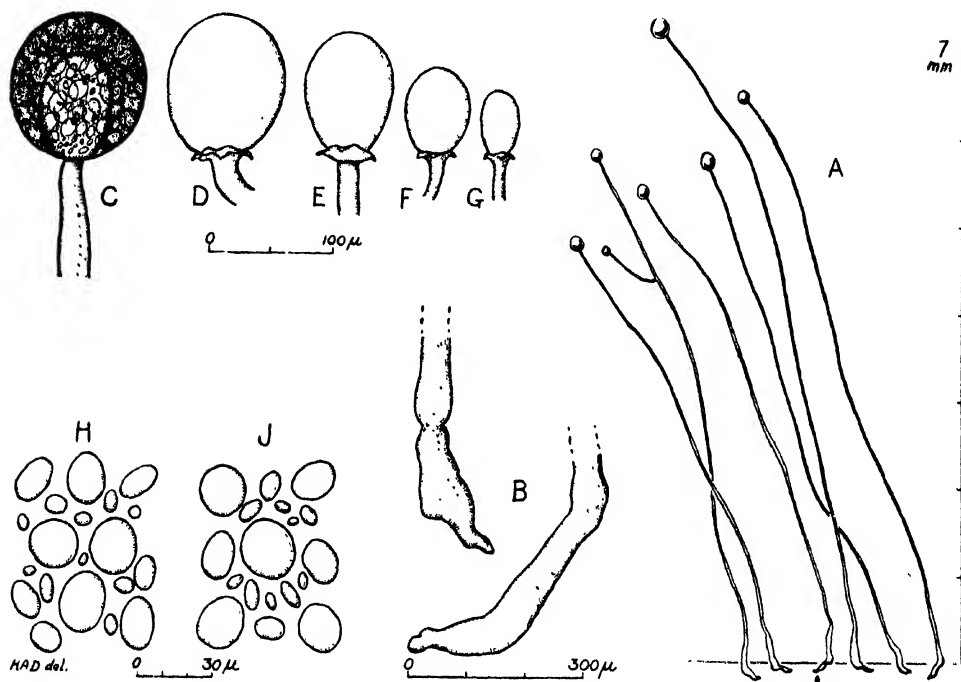


Fig. 3. *Mucor inaequisporus* Dade. A, habit, $\times 10$ d.; B, bases of sporangiophores, $\times 78$ d.; C, a mature sporangium, $\times 165$ d.; D-G, columellae, $\times 165$ d.; H, J, spores from vegetations on hog-plum fruit and potato-dextrose agar respectively, $\times 330$ d.

trose agar, when growth is unrestricted on plates covered with bell-jars, they attain a height of 4 cm.; their diameter is $35-49\mu$, and they have broad short bases $40-70\mu$ diam.; they have orange-coloured contents. The sporangia are spherical, averaging $100-150\mu$ diam. on the fruit, maximum 200μ ; on potato-dextrose agar the average diameter is 100μ , maximum 150μ ; they are golden brown when young, dark brown when mature, by reflected light; the membrane is diffuent, and has no investment of crystals. The columellae are pyriform, subspherical in large individuals, ranging from 40×30 to $100 \times 90\mu$, average about $80 \times 60\mu$, with rather deep brownish orange contents.

The spores are exceedingly variable in size (the outstanding characteristic of this species), 3.5×2.5 to $25 \times 20\mu$, the smallest being elliptic, sometimes reniform, the largest nearly round, the intermediate sizes broadly elliptic; spores of all sizes come from each individual sporangium; they are pale yellowish and are not mingled with oil droplets.

The gross colour of vegetations is similar in culture on potato-dextrose and corn-meal agar and on hog-plum fruits, and strongly resembles the colour of the latter: Ridgway IV.19.—apricot to IV.19.b light cadmium-yellow. The fungus grows more freely on potato-dextrose than on corn-meal agar.

The long aerial sporangiophores are strongly heliotropic.

TAXONOMY

This species is not closely comparable with any other previously described. In spite of the occasional side branches, it is as typically a *Monomucor* as is *Mucor Mucedo*, which also produces lateral branches occasionally. In other features it also resembles *M. Mucedo*, to which it is evidently more nearly related than to other species; but it differs from *M. Mucedo* markedly in the colour of the sporangia and sporangiophores, in the form of the columella, the absence of crystals on the membrane, the more extreme variation in spore size, and the different shape of the spores.

These differences are considered to justify the proposal of a new species, *M. inaequisporus*.

DIAGNOSIS

Mucor inaequisporus n.sp.

Hyphae fertiles erectae sed heliotropicae, typice non ramosae, aliquando cum ramulis singulis lateralibus, hyalinae sed materiam aurantiacam continent; in fructibus hospitis 0.5–0.75 cm., in agaro ad 4 cm. long., $35\text{--}49\mu$ crass., basibus brevibus latisque $40\text{--}70\mu$ crass.

Sporangia globosa, in juventate mellea, in maturitate badia, cum membrana diffuente, crystallis carentia; in fructibus plerumque $100\text{--}150\mu$, ad 200μ diam., in agaro plerumque 100μ , ad 150μ diam.

Columellae piriformes vel si magnae subglobosae, hyalinae sed materiam aurantiacam continent; $30\text{--}40 \times 90\text{--}100\mu$, plerumque $60 \times 80\mu$.

Sporae in sporangiis singulis magnitudine dissimiles, plerumque ellipsoideae vel magnae subglobosae vel parvae reniformes, subflavidae, guttulis olei carentes, $3.5 \times 2.5\text{--}25 \times 20\mu$, plerumque $20 \times 15\mu$.

Cultus in agaro armeniacus.

In fructibus maturis abstrictisque *Spondiadis monbin* L. in Gold Coast, Africae Occidentalis.

ABSIDIA CRISTATA N.SP. (Gold Coast No. 554)

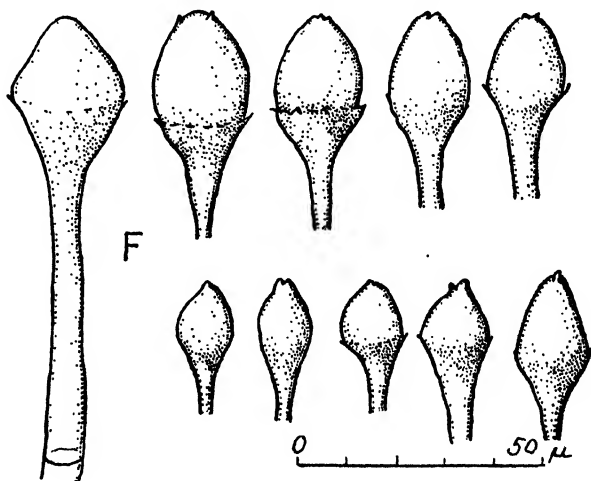
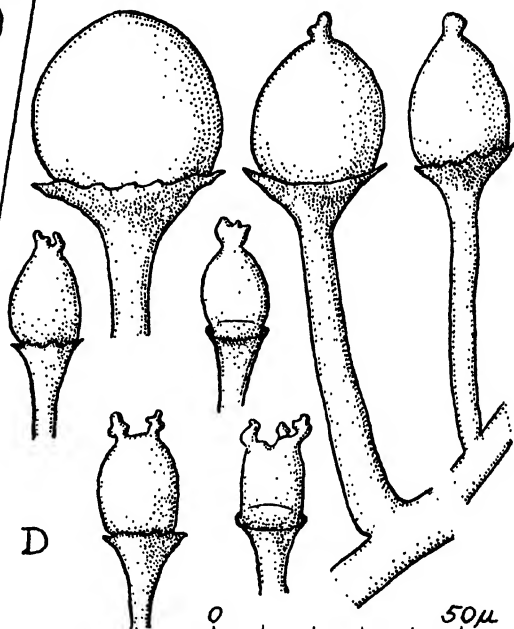
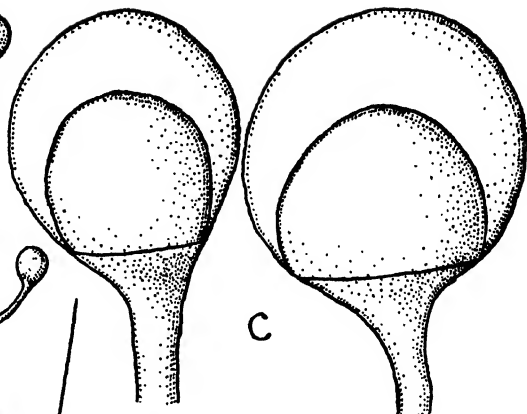
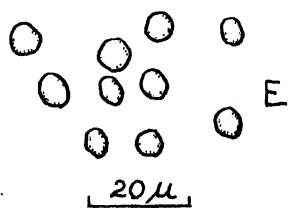
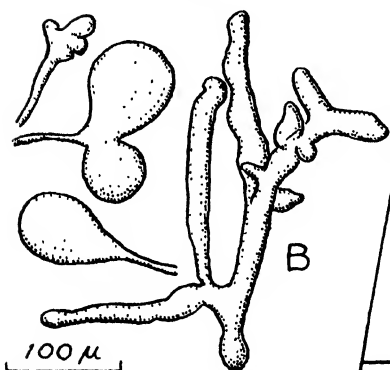
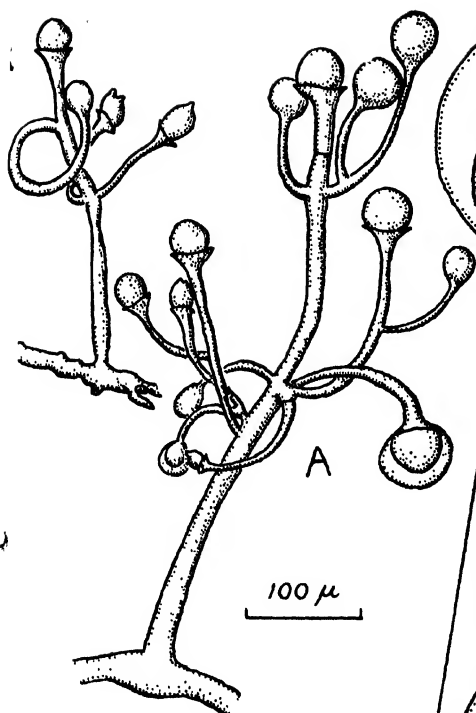
This fungus was isolated from cassava stems on which it was saprophytic following anthracnose, and from fermented cacao on which it is an uncommon mould. In culture it is characterized by two distinct types of growth—it first forms a low sward of short, rather thickly branched sporangiophores, and later long cobwebby sporangiophores with similar racemose branching, but with the laterals at much longer intervals. It was at first thought that two fungi were involved, but observation in culture showed that the two forms belong to the same species.

The fertile hyphae are hyaline, up to 20μ thick, the lateral branches narrower, up to $8-9\mu$ thick. The sporangia are nearly spherical, but with the infundibuliform apophysis typical of *Absidia*, and dark brown to black in appearance. The columellae are spherical to oval, of diverse form: in terminal sporangia they are large, globose, up to 40μ diam., smooth on the summit; in smaller lateral sporangia they are oval, with a distinctly formed papilla on the summit; while in the smallest laterals they are narrowly oval, nearly twice as long as broad, with one or a few apical processes which may be simple papillae but are more often highly developed outgrowths of more or less complex form; all columellae are fuliginous, the colour extending into their infundibuliform bases. The spores are irregularly rounded, with a tendency to polygonal outline, $4-7\mu$ diam. and smoky brown; there is a suggestion of punctate markings. The zygospores, chlamydospores and gemmae have not been seen. In culture abnormal growths occur, taking the form of giant hyphae and of large tuberculate developments of sporangial heads, all with dense granular contents.

Of the species of *Absidia* previously described, *A. Lichtheimii* (Lucet & Cost.) Lendner(?) seems to be nearest to the present fungus. An authentic culture of *A. Lichtheimii* was provided by the Imperial Mycological Institute for comparison. This produces smaller colourless spores $2-3\mu$ diam., the columellae are more ovoid in form, and the apical processes are only very slightly developed. It is difficult to find any close resemblance between the two fungi, and it therefore seems proper to describe the Gold Coast fungus as a new species, with the name *A. cristata*.

Legend to Fig. 4.

Fig. 4. *Absidia cristata* Dade. (A-D, *A. cristata*; F-G, *A. Lichtheimii* for comparison.) A, habit, compact type, $\times 156$ d.; B, abortions produced in culture, $\times 156$ d.; C, large terminal sporangia, $\times 660$ d.; D, a series of columellae, the largest from a terminal sporangium, the others from smaller lateral sporangia, $\times 660$ d.; E, spores, $\times 660$ d.; F, a series of columellae of *A. Lichtheimii*, $\times 660$ d., for comparison with D; G, spores of *A. Lichtheimii*, $\times 660$ d., for comparison with E.



H.A.D. del.

DIAGNOSIS

Absidia cristata n.sp.

Hyphae fertiles hyalinae racemosae biformes: 1. breves dense racemosae; 2. elongatae; sporangia magna terminales et minores in ramis lateralibus gerentes. Primae stirpes ad 20μ , rami laterales ad $8-9\mu$ crass.

Sporangia pyriformia vel subglobosa, ad 60μ diam., apophyse infundibuliformi, in maturitate badia vel subnigra.

Columellae subglobosae vel ovaes; in sporangiis terminalibus magnae, ad 40μ crass., cum apicibus levibus; in sporangiis lateralibus minoribus ovaes, semper cum apicibus papillatis; in sporangiis lateralibus minimis subanguste ovaes, singulos vel paucos, simplices vel saepius magnas multiformes, apicales processus gerentes; fuliginosae.

Sporae incondite globosae vel ovaes, subpolygoniae, aliquando paululum punctulatae, fumosae-fuliginosae, $4-7\mu$ diam.

Zygosporae ignotae.

Saprophila in stirpibus *Manihotis utilissimae* Pohl. et in seminibus *Theobromae Cacao* L., in Gold Coast, Africae Occidentalis.

The author is indebted to Messrs S. F. Ashby, E. W. Mason and R. H. Bunting for assistance in consulting literature not available in Africa, and for their opinions on diagnoses.

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A PYTHIUM WILT OF PRIMULA CAUSED BY *PYTHIUM SPINOSUM*, SAWADA

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W. B. COLLINS, B.Sc.

University College of South Wales and Monmouthshire

(With 7 Text-figures)

INTRODUCTION

IN October 1934 a serious epidemic disease occurred in the experimental plants of *Primula sinensis* growing at the John Innes Horticultural Institution. The seedlings wilted in the seed pans, causing considerable mortality among the many hybrid varieties grown for experimental purposes. A preliminary examination of the seedlings was made by Dr G. H. Pethybridge, who came to the conclusion that the epidemic was due to the parasitic fungus, apparently a member of the Pythiaceae, which he found growing in the roots. Some of the material was sent to the senior author with a request that he would make a more detailed investigation of the fungus and determine its identity if possible.

Meanwhile, investigations carried out by Miss D. M. Cayley at the John Innes Horticultural Institution showed that, although finally the seedlings became infected by a fungus, the immediate cause of the disease was the soil in which the *Primula* plants were growing. This soil was sterilized for forty minutes at 100° C. and then left for about six weeks in a soil bin outside before it was used for potting the seedlings. Miss Cayley's work indicated that heating the soil broke down the humus of the compost, rendering the soil too dense and thus preventing normal growth of plants in it. "The main cause of the apparent epidemic of wilt is that the roots of the plants are not keeping pace with the top growth, and as they grow larger the excessive evaporation makes them wilt and collapse and thus become an easy prey to secondary organisms".* Miss Cayley tried the effect of repotting seedlings which were attacked, in smaller pots in a much more porous soil, with the result that, within a few weeks, the plants recovered. Similar results were obtained with infected plants which had been sent to the authors and which were treated in the same way in the experimental greenhouse at Cardiff. In such plants the spread of the fungus is arrested.

It was felt, however, that it was still desirable to determine the nature of the fungus in the roots, since a preliminary study had failed to identify it.

* In a letter to the senior author.

METHODS

The material received consisted of fresh seed and infected seedlings, some of them still in their original pots of infected soil. Some of the seedlings which were planted in infected soil have remained infected though they have not died. Others, which were repotted into the sterile soil used at Cardiff, grew healthily, and all signs of the fungus disappeared. Seed sown, both in the sterile soil and also on agar plates, never contracted the fungus, neither was any fungus isolated from the agar itself, although it was found that the infecting fungus would thrive on the medium. From this it was concluded that the fungus was present in the soil and did not come from the seed of *Primula sinensis*.

The fungus was isolated from infected roots by washing them in mercuric chloride and then teasing out the inner parts of the roots into sterile distilled water contained in small glass jars. After a few days growth appeared on pieces of hemp seed which had been introduced into the jars and the fungus passed through its entire life history at laboratory temperature under these conditions. Several separate isolations were made and the same organism was always isolated.

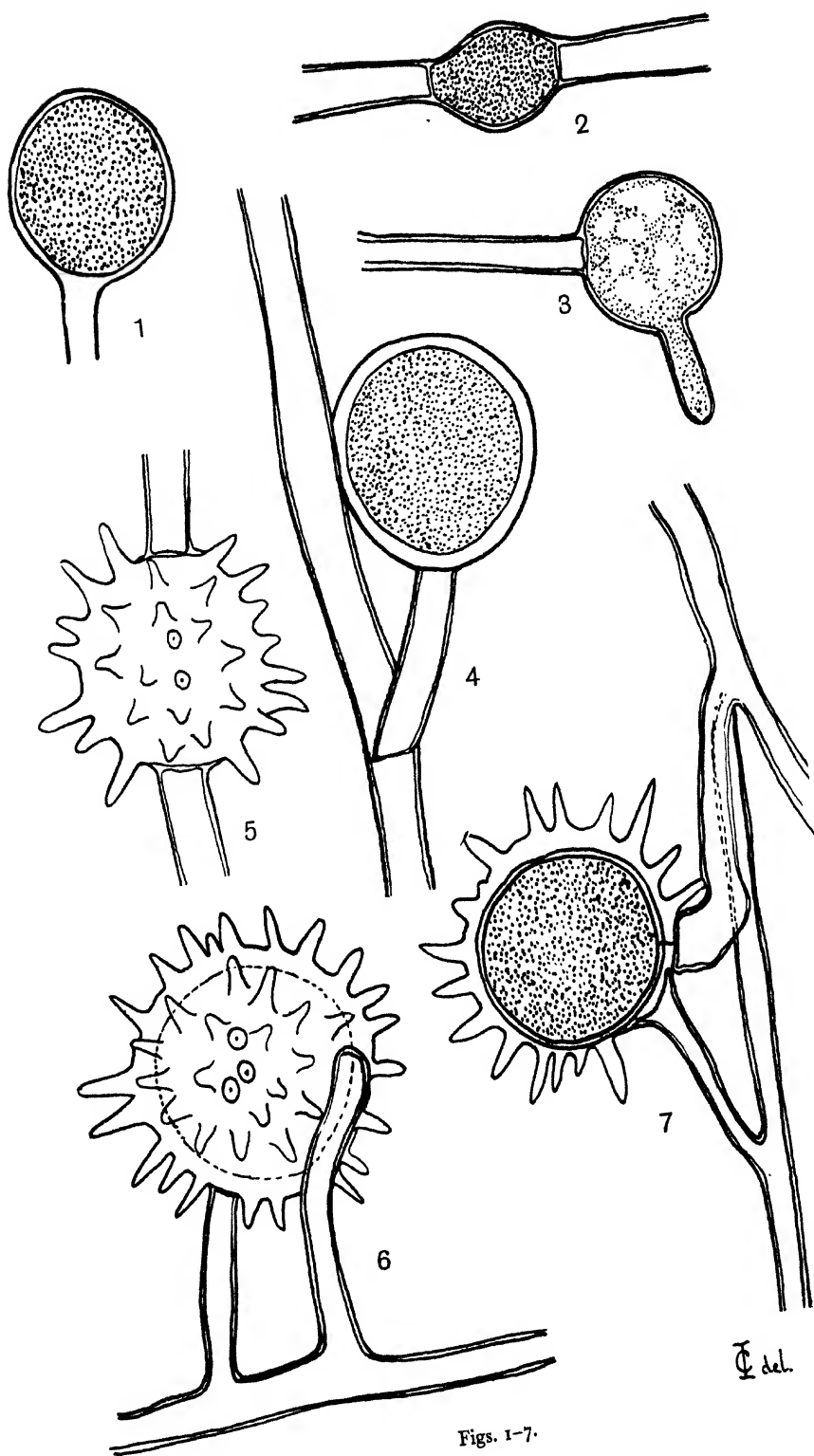
LIFE HISTORY OF THE FUNGUS

The organism develops a fine mycelium forming a rather tufted growth on the hemp seed. The delicate hyphae measure $2-5\mu$ diam. and, when young, are devoid of transverse septa. Septa appear in older hyphae, and in old mycelia regular septation is present. After a short time in culture the fungus begins to reproduce asexually by means of conidia. No zoosporangia are formed and zoospores are never differentiated. The conidia form abundantly in culture and usually develop terminally (Figs. 1—4), though intercalary conidia

Legend to Figs. 1-7.

The drawings were made with a camera lucida at table level using a Zeiss 4 mm. objective and compensating ocular $\times 20$; tube length of 160 mm. giving a magnification of 1580. Reproduction is without reduction.

- Fig. 1. A single apical conidium terminating a main hypha.
- Fig. 2. An intercalary conidium at a young stage in development.
- Fig. 3. A terminal conidium germinating by means of a single germ-tube while still attached to the parent hypha.
- Fig. 4. A conidium developing on a lateral branch hypha, which has been cut off by a septum: the type of development on an older mycelium.
- Fig. 5. An intercalary oogonium at a young stage in development; no antheridium was present at this stage.
- Fig. 6. A mature oogonium and antheridium both arising from the same main hypha, but the antheridium not springing from the oogonium stalk. The dotted line indicates the position of the oosphere.
- Fig. 7. A mature oogonium and antheridium after discharge of the contents of the latter into the oogonium. The drawing shows the appearance in median section with the oospore delimited. The antheridial stalk springs from a triple branch of the main hypha.



♂ del.

Figs. 1-7.

appear rarely. The wall of the conidium is usually smooth, though sometimes short spines may be developed. The conidia are roughly spherical and measure $16.5\text{--}34.3\mu$ in diameter (average 26.4μ).^{*} They germinate by means of a single germ-tube (Fig. 3) and never by the production of zoospores.

In older cultures sexual reproduction occurs. The antheridia are stalked, androgynous, and rarely arise from the oogonial stalk. The oogonia, which are usually formed terminally, are spherical, measuring $16.0\text{--}28.4\mu$ (average 19.0μ), and densely covered with spines from $3.5\text{--}8\mu$ long (Figs. 5-7). Each oogonium, after fertilization, gives rise to a single oospore which usually completely fills the oogonium. It is spherical and measures $13.0\text{--}20.3\mu$ diam. (average 15.4μ). Its wall is smooth. Germination of the oospore has not been observed.

DISCUSSION

The fungus is a member of the genus *Pythium*, most closely agreeing with *P. spinosum*, a culture of which was obtained from the Centraalbureau voor Schimmelcultures: the following table shows the comparative measurements:

	<i>Pythium spinosum</i> Sawada	Fungus isolated from seedlings
Hyphae	2-5 μ diam.	2-5 μ diam.
Conidia	14-33 μ (av. 22.4 μ) diam.	16.5-34.3 μ (av. 26.4 μ) diam.
Oogonia	17-24 μ (av. 19.7 μ) diam.	16.0-28.4 μ (av. 19.0 μ) diam.
Spines	5-8 μ long	3.5-8 μ long
Oospores	12-21 μ (av. 16.6 μ) diam.	13.0-20.3 μ (av. 15.4 μ) diam.

These comparisons show that the fungus which attacked the seedlings was *Pythium spinosum* Sawada. This species was first described by Sawada & Chen (3) as a parasite of *Antirrhinum* seedlings in Formosa. They showed by inoculation experiments that the fungus could infect young plants of a number of other common species, and its ability to attack the seedlings of *Primula sinensis* is in no way remarkable. The important question, however, is whether this species has been introduced into this country from Japan. We are informed that records from the John Innes Horticultural Institution show that no plants had been introduced from Japan during a period which would make infection probable. It seems likely therefore that the species occurs wild in this country, but that it is only by chance that its presence has become recognized. The genus *Pythium* has as yet been only superficially investigated in Great Britain, and experiments carried out in the Department of Botany at Cardiff indicate that the number of common species inhabiting the soil is far greater than is generally recognized, and that a number of species apparently not so far recorded for this country have been found.

* Each average is based upon twenty-five measurements.

The occurrence of this species as an epidemic parasite of *Primula* seedlings, however, seemed sufficient to justify special notice being taken of the fungus.

Specimens of *Pythium spinosum* Sawada, both from the Baarn culture and also from the isolation on Primulas, have been preserved in the senior author's herbarium.

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NOTES ON ENTOMOGENOUS FUNGI

By T. PETCH

101. *ENTOMOPHTHORA PLANCHONIANA* Cornu

CORNU, in *Bull. Soc. Bot. France*, xx (1873), 189, 190, gave an account of an *Entomophthora*, found on an aphid on elder at Montpellier, for which he proposed the name *E. Planchoniana*, because apparently the same fungus had been observed previously by Planchon on an aphid on a vetch. No formal description was given, and the following details give all the relevant information.

The affected insects were brick-red, instead of the normal black, and their abdomens were enormously swollen. The sporangia were globose, apiculate, filled with a refringent plasma, and containing in the centre a spore of the shape of a "toupie d'Allemagne". The fungus resembled *Empusa Muscae*, but could not be referred to that species. In a moist atmosphere the sporangia produced secondary sporangia, similar to the primary. Under special conditions of humidity the wall of the sporangium remained adherent to the wall of the spore, and the shape of the whole was then ovoid. In a subsequent paper by Brongniart & Cornu (*Ass. Franç. Avanc. Sci.* Paris, 1878), which deals with an attack of an *Entomophthora* on syrphids, there is a footnote, stating that according to the degree of humidity *E. Planchoniana* produces ovoid-oblong spores without sporangia, or sporangia soldered to the spore.

By "sporangia" Cornu meant what are now known as conidia. The true conidium of an *Entomophthora* is formed within a mother cell, the wall of which is normally in close apposition to that of the conidium, so that the whole appears to be a single spore. But in some species, with excess of moisture, the wall of the mother cell separates from the conidium proper, so that the latter floats free in a large spherical cell. As regards the shape, "toupie d'Allemagne", according to the dictionaries, means a humming top. But in the second note referred to above the same term is applied to the conidium of an *Entomophthora* which was undoubtedly *Empusa Muscae*. It is clear, therefore, that *Entomophthora Planchoniana* had conidia which resembled those of *Empusa Muscae*, and under suitable conditions of humidity the outer wall separated from the true conidium, leaving the latter floating in a large spherical cell.

Thaxter, in *Entomophthorae of the United States* (1888), applied Cornu's name to an American species "with much hesitation". He pointed out the discrepancies between Cornu's two descriptions of the spore, but apparently did not realize that Cornu's "toupie d'Alle-

magne" was the same shape as the spore of *Empusa Muscae*. Thaxter's American species had globose spores, and he stated that he had never seen a separation of the outer wall in spores of that type.

In September 1933, I found a single specimen of an *Empusa* on a detached aphid in Deepdale, Barnard Castle, which resembled *E. Muscae*, and was recorded under that name. Subsequently, I received specimens of the same fungus on an aphid on raspberry from Dr H. S. Fawcett, collected at Puyallup, Washington, U.S.A., June 1934; and in May 1935 I found it in quantity on an aphid on green-gage in my garden at North Wootton.

The insect is fastened to the leaf by short rhizoids from the under side of the body, which are not visible until it is forcibly detached. The conidiophores cover the insect either in a continuous even stratum or in irregular confluent masses. The insect is red, and the layer of conidiophores is pale reddish. The conidiophores are simple and there are no cystidia. The conidia are ejected with a drop of plasma, as in *E. Muscae*, are bell-shaped or globose with a truncate base, and are furnished with an apical point; the sides are turned outwards slightly towards the base, which is slightly convex; they measure $17-23 \times 12-20\mu$. In a spore print in a damp chamber, the outer wall frequently separates from the conidium proper, so that the latter floats free in a spherical cell about 30μ in diameter. The secondary conidia are the same shape as the primary, with or without an apical point, but smaller, $15-20 \times 12-14\mu$, and are produced by direct budding from the primary conidia. The spore is shaped like an old-fashioned German spiked helmet.

This species appears to fit Cornu's description better than that for which the name was adopted by Thaxter. It resembles *E. Muscae*, but has smaller conidia, and the insect is attached to the plant by rhizoids. In spite of the latter feature, however, it must be referred to *Empusa*, not *Entomophthora*, as the conidiophores are simple. I propose to adopt Cornu's name for this species, viz. *Empusa Planchoniana* (Cornu) Petch non Thaxter, while the species for which the name was employed by Thaxter may be known as *E. Thaxteriana*.

In *Ann. Mag. Nat. Hist. Ser. 5*, xviii (1886), 4, pl. 3, figs. 1-13, Phillips recorded *Entomophthora ferruginea* on *Aphis rumicis*. The fungus had been compared with *Entomophthora Aphidis*, and found to be different. The details given are not sufficient for an accurate determination, and the drawings are equally inadequate, but some of the latter show narrow bell-shaped conidia, sometimes with an apical point, and could be matched in a squash preparation of *Empusa Planchoniana*. *Entomophthora ferruginea* would appear to agree better with *Empusa Planchoniana* than with *Entomophthora Aphidis*, to which Thaxter referred it. Unfortunately, no specimen has been found in the Phillips collection in Herb. British Museum.

The synonymy of the two species considered above is:

Empusa Planchoniana (Cornu) Petch; *Entomophthora Planchoniana* Cornu in *Bull. Soc. Bot. France*, xx (1873), 189; *Entomophthora ferruginea* Phillips in *Ann. Mag. Nat. Hist.* ser. 5, xviii (1886), 4; non *Empusa Planchoniana* Thaxt., *Entomophthorae of the United States* (1888), p. 165.

Empusa Thaxteriana Petch, nom.nov.; *Empusa Planchoniana* Thaxter, *Entomophthorae of the United States* (1888), p. 165.

102. *ENTOMOPHTHORA PYRALIDARUM* Petch, n.sp.

In December 1923, large numbers of "grass moths" (Pyralidae), which had been attacked by an *Entomophthora*, were found in the Arboretum, Royal Botanic Gardens, Peradeniya, Ceylon, attached to the trunks of trees. A similar occurrence was observed at Vavuniya, in the low-country, in the same month; and in the following month examples were found at Kandy. Specimens were submitted to Prof. Thaxter, who stated that he was not acquainted with the fungus.

The moths were attached to the bark by numerous rhizoids, which were simple, sometimes expanding towards the extremity, and arose chiefly from the abdomen. The conidiophores are branched, and form a loose stroma, principally over the abdomen, becoming aggregated in small tufts. The conidia are pyriform, or oval with a conical or obtuse papilla, $16-28 \times 12-16\mu$, or globose, $18-28\mu$ diameter, with a conical papilla, $4-8\mu$ high, not apiculate. Oblong, thick-walled chlamydospores, with a large central gutta, $22-28 \times 13-14\mu$, occur terminally on the hyphae. Internally, zygospores (?), spherical, smooth, hyaline, $14-24\mu$ diameter, were found. The contents of the conidia are uniformly granular.

Entomophthora pyralidarum Petch, n.sp.—Hyphis rhizoideis simplicibus, interdum apicem versus incrassatis, praecipue ex abdomine oriundis; conidiophoris ramosis praecipue abdomine insessis, fasciculatis; conidiis pyriformibus vel ovalibus, conice vel obtuse papillatis, $16-28 \times 12-16\mu$, vel globosis, $18-28\mu$ diam., papilla conica, $4-8\mu$ alt., instructis; chlamydosporis oblongis, episporio crasso, terminalibus, $22-28 \times 13-14\mu$, gutta magna centrali; zygosporis (?) internis, sphaericis, levibus, hyalinis, $14-24\mu$ diam. In Pyralidis, Ceylon.

103. *HYPOCRELLA CORNEA* Petch, n.sp.

Five gatherings of this species, on an Aleyrodid on *Rubus*, were included in a collection of entomogenous fungi from Kwangsi Province, China. In general appearance it resembles *Hypocrella Mollii*, but differs from that species in texture. It is white or yellowish, but becomes amber-coloured when rubbed, having a cortical zone which is hard and horny when dry. There are indications that it may be viscid when fresh. The ostiola are dark amber, not brown as in *H. Mollii*.

Hypocrella cornea Petch, n.sp.—Stromatibus usque 2.5 mm. diam., orbicularibus, discoideis vel depresso-pulvinatis, aequalibus vel leniter tuberculatis, albis vel flavescentibus, pruinosis, cortice sicco duro corneo, medio albis, laxis; peritheciis immersis, praecipue ad marginem, late ampullaceis, 0.3 mm. alt., 0.2–0.25 mm. diam., pariete hyalino vel flavo; ascis cylindraceis vel fusoides, $130\text{--}200 \times 8\text{--}10\mu$; articulis sporarum cylindraceis, dein ovalibus, truncatis, $7\text{--}9 \times 1.5\text{--}2.5\mu$. On an Aleyrodid on *Rubus*, Kwangsi Province, China.

104. *CORDYCEPS FORMICIVORA* Schroet.

In *Krypt.-Flora von Schlesien*, III, 2 (1894), 276, Schroeter described *Torrubia formicivora*, which had occurred on *Formica ligniperda* in Silesia. The stroma consisted of a very short stalk, 1–2 mm. long, 0.5 mm. diameter, sometimes almost lacking, and a globose or hemispherical head, up to 2 mm. diameter, blackish, brownish internally. The perithecia were crowded, oval, up to 0.3 mm. diameter, semi-immersed, with a conical ostium. The asci were cylindric, generally attenuated below, $150\text{--}170 \times 6\text{--}7\mu$, eight-spored, the ascospores being linear, $100\text{--}110\mu$ long, $2.5\text{--}3\mu$ broad, at first continuous, then septate, hyaline or pale yellow. The conidial clava was awl-like, simple, or divided above with diverging branches, up to 3 mm. high, 0.5 mm. diameter, blackish brown, generally pale violet-brown at the apex. Schroeter's fungus was clearly not *Cordyceps myrmecophila* Ces., and his description did not suggest any previously known species.

In *Proc. Amer. Phil. Soc.* LXXIV (1934), 263, Prof. E. B. Mains recorded *C. formicivora* Schroet., noting: "This curious species has been reported by Povah. He obtained one specimen on an ant at Rock River, Michigan, in 1927. This is apparently the first report for North America. Povah, however, states that Prof. Thaxter had collected it on several occasions at York, Maine."

I have examined specimens collected by Thaxter at Kittery Point, Maine, August 1892 (Farlow Herb. 6149), and at York, Maine (Farlow Herb. 6154), all on ants.

In No. 6149, the ants are fastened to the substratum by brown mycelium, with rather loose brown mycelium along the sutures, legs, etc. One has two short brown clavae arising from behind the head, each bearing a black, verrucose, lateral, perithecial plate. The other has three adjacent perithecial plates behind the head, but the clavae are so short that the plates appear sessile.

In No. 6154, one specimen has two shortly stalked perithecial plates behind the head. The other bears two clavae, one from behind the head, 3 mm. high, with two lateral perithecial plates at the base, and conidia on the upper linear part of the clava, and the other, conidial only, arising from a leg. The conidial clavae bear *Hirsutella* conidio-phores.

Thaxter assigned these specimens to *Cordyceps unilateralis*, but subsequently crossed out that name. But they are, I think, that species, though smaller than is usual in the tropics. There is another North American specimen of *Ophiocordyceps unilateralis* in the Thaxter collection, Cullowhee, N.C., 15 June 1896 (Farlow Herb. 6161).

Rereading Schroeter's description in the light of these specimens, it seems clear that he was dealing with a small example of *Ophiocordyceps unilateralis*. The American specimens have been correctly assigned to Schroeter's species, but his name must be regarded as a synonym. His record appears to be the only reported occurrence of this species in Europe.

105. *CORDYCEPS SUPERFICIALIS* (Peck) Sacc.

In the 28th Report *N.Y. State Museum*, p. 70, Peck described *Torrubia superficialis*, which he stated was intermediate between *Cordyceps acicularis* and *C. Ravenelii*. As these two latter are the same species, an *Ophiocordyceps*, Peck's name was reduced to a synonym in *Trans. Brit. Mycol. Soc.* xviii (1933), 60.

Peck's type has now been examined by Prof. E. B. Mains, who has published the following redescription of it in *Proc. Amer. Phil. Soc.* lxxiv (1934), 268.

"Stromata 'vinaceous-buff', 2.5 and 3.5 cm. long, 1.5-2 mm. in diameter, the stipes 10 and 15 mm. long, fertile portions 15 mm., one with sterile acuminate apex, 5 mm. long, the other with apex apparently broken off; perithecia superficial, free, somewhat crowded, 'army-brown', ovoid, 456-660 \times 450-564 μ , some with apex collapsed or broken off; asci cylindric, 170-230 \times 7-9 μ ; ascospores hyaline, filiform, probably about equalling the asci in length, breaking into segments 14-30 \times 1.5-2 μ ."

In the Farlow Herbarium there is a specimen, No. 6130, collected by Thaxter, "on *Melolonthae* (?) in mossy woods, West Haven, Conn., 26 July 1889", which is apparently referable to Peck's species. The clavae are up to 2.5 cm. high, with a flexuose, longitudinally grooved, smooth, terete, brown stalk, 0.5 mm. diameter. The fertile portion of the clava is about 1 cm. long, 1 mm. diameter, cylindric, dark brown, almost black. Sometimes the stem is continued above as a sterile yellow apex. The perithecia are superficial, crowded, ovate, 0.36-0.4 mm. high, 0.3-0.36 mm. diameter, dark brown, brown pruinose, apex obtuse. The asci are subcylindric or narrow-clavate, 150-175 μ long, 7 μ diameter, and the free part-spores measure 16-24 \times 1.5 μ . By transmitted light, the wall of the perithecium is rather dark brown. The host appears to be a beetle larva, but not a *Melolonthid*.

I have examined another specimen, collected at Norwood, Mass., 27 August 1933, on larvae buried about one-half to one inch deep in

leaf mould and soil in woods composed chiefly of *Acer rubrum*. In this, the part-spores were cylindric, sometimes attenuated to one end, $14-27 \times 1.5-2\mu$.

This is a *Cordyceps*, not *Ophiocordyceps*, differing from *O. acicularis* in the shape of the perithecia and the character of the ascospores. The length of the part-spores is exceptional for an entomogenous *Cordyceps*.

106. *CORDYCEPS SUBSESSILIS* Petch, n.sp.

It is doubtful whether this species should be referred to *Cordyceps* or to *Torrubiella*. It is represented in the Farlow Herbarium, *ex* Herb. Thaxter, by two specimens, both on a coleopterous larva in wood, No. 6145, Burbank, Tennessee, August 1896, and No. 6155, Cranberry, North Carolina, 1887.

In No. 6145, the fungus has an irregular, strap-shaped pseudostalk, 5 mm. high, 1 mm. broad, smooth, yellowish, with brown, horny patches here and there, which bears a cluster, 2 mm. diameter, of perithecia at the apex. In No. 6155, the larva has apparently been cut longitudinally and the specimen broken up; all the perithecia are now detached, and there are fragments of a thin, brown, horny clava. Judging from these specimens, it would appear that the fungus produces perithecia as soon as its mycelium reaches the surface of the wood, the apparent stalk being merely the strand of mycelium in the wood, probably in the insect bore-hole.

The perithecia are remarkably large, 1-1.5 mm. high, 0.33-0.44 mm. diameter. They are numerous, superficial, free, crowded, narrow flask-shaped, collapsing laterally, amber, glabrous, or with slight, white or yellowish mycelium between their apices. The asci are cylindric, capitate, $240 \times 5\mu$, and the part-spores cylindric, $3-7 \times 1\mu$.

Cordyceps subsessilis Petch, n.sp.—Stipitibus irregularibus, flavis dein brunneis, ligno immersis, apice perithecia conferta ferentibus; peritheciis liberis, anguste ampullaceis, 1-1.5 mm. alt., 0.33-0.44 mm. diam., a latere collabascentibus, succineis, glabris; ascis cylindraceis, capitatis, $240 \times 5\mu$; articulis ascosporarum cylindraceis, $3-7 \times 1\mu$. On larvae of Coleoptera in wood, North America.

107. *CORDYCEPS CURCULIONUM* (Tul.) Sacc.

Cordyceps curculionum was figured by Robin as *C. entomorrhiza*, and his mistaken identification was corrected by Tulasne. It appears to have been fairly well known in the middle of the last century, but it is not well represented in herbaria at the present day. Cooke, writing of this species in his *Vegetable Wasps and Plant Worms*, stated: "It is generally known amongst entomologists that specimens of *Curculio* are collected in tropical countries, and sent home with the collections, which have club-shaped parasites attached to them. Foreign col-

lectors, or collectors abroad, have generally a keen eye to the monetary value of such 'varieties' and are seldom satisfied with their weight in gold to part with them. The golden opportunities are perhaps rather too rare." Whatever the reason, there are few specimens in herbaria. Incidentally it may be noted that their weight in gold would not be of great value, even at current rates. The object of the present note is to call attention to a feature, probably related to *Cordyceps curculionum*, which does not appear to have been described.

In Herb. British Museum there is a specimen of a fungus on a weevil (*Helipus* sp.), collected by Bates at Tapajos, Brazil (B.M. 165). The fungus emerges from the insect along the sutures, in the form of pale brown, loose, conical tufts, up to 1.5 mm. high, 0.6 mm. diameter at the base, tapering to a more or less acute apex. These are united at their bases, or laterally coalescent in an irregular fringe. The tufts are composed of loosely interwoven hyphae, 3–4 μ diameter, smooth and flexuose. There is no central core of agglutinated hyphae, and no conidiophores or conidia have been found on them. Another specimen in Herb. British Museum (No. 166), also on a weevil (*Sternechus* sp.), collected by Bates at Santarem, Brazil, is similar to the foregoing, while in Herb. Berlin there is another example, collected by Tessmann (No. 83) in Peru.

None of these specimens could be assigned to any species on their own evidence. There is, however, another specimen in Herb. British Museum (No. 175), on a weevil (*Helipus* sp.), from Guayaquil, 26 July 1871, which, in addition to these growths along the sutures, bears a perithecial clava of *Cordyceps curculionum*. It would appear, therefore, that these sterile growths are related to that species. They are not the conidial stage, which is either a *Hirsutella* or a *Hymenostilbe*, but this has not yet been seen in a fit condition for determination.

108. *CORDYCEPS EROTYLI* Petch, n.sp.

In *Vegetable Wasps and Plant Worms*, p. 115, Cooke, writing of *Cordyceps curculionum*, stated: "It is not improbable that the coleopterous insect alluded to by Mr Gray as belonging to the family Erotylidae (the *Erotylus taeniatus* of Columbia) was attacked by a form of this same *Cordyceps*." He then quoted: "It is remarkable for producing occasionally from the head, from between the head and thorax, and from the sides of the body, a number of very slender vegetable appendages, the apex of each ending in a small tuberculated head of a yellowish colour."

There are two specimens of a *Cordyceps* on *Erotylus* sp. in the Farlow Herbarium. No. 2625 was collected by Thaxter at Verdant Vale, Trinidad, 1912–13. The fungus is on the mature beetle, lying in its burrow in wood, and the clavae emerge from the entrance to the

burrow. The insect is sparsely covered with yellowish mycelium, and the numerous clavae arise from the sutures on the upper and under surface, sometimes most from the head, sometimes most from the hinder end. The clavae are up to 15 mm. high, simple, or bifurcating above. The stalk is 0.15–0.25 mm. diameter, externally yellowish pruinose, internally subtranslucent and red-brown. The head is clavate or subcylindric, at first yellowish and sterile. The perithecia in an early stage are immersed, or partly immersed, in yellowish, loose tissue, but become superficial, amber, then brownish red, sometimes confined to one side of the head. The head is 1.5–4.5 mm. long, 0.36–0.75 mm. diameter, and the perithecia are conoid to flask-shaped, 0.4 mm. high, 0.18 mm. diameter, slightly tomentose. The asci are about 250 μ long, 4 μ diameter, cylindric, capitate, and the ascospores linear, 0.75 μ diameter. Part-spores were not found.

Another specimen, No. 6120, Verdant Vale, Arima, Trinidad, 1913, bears two clavae like those of No. 2625. In addition, delicate strands of mycelium, white becoming brown, extend from the insect along the sides of the boring to the entrance, and thence radiate over the surface of the wood in a patch about 2 cm. diameter. These strands bear *Spicaria* conidiophores, simple, short, about 130 μ high, with a stout septate stem, 6 μ diameter, and a head, 25–45 μ high, consisting of two or three whorls of flask-shaped phialides, 6–14 \times 2–2.5 μ . The conidia are cylindric, ends rounded, 3–5 \times 1–1.5 μ .

A third specimen, Farlow Herb. 2582, collected by Thaxter in the same locality, December 1912, is marked by him, "*Isaria* of *Cordyceps* on *Erotylid*; was with the *Cordyceps* on the same host." A white mycelium emerges along the sutures and spreads in strands along the legs, passing over to the wood, on which it forms a white sheet. Conidiophores occur on the mycelium and on minute, white, loose clavae up to 1.5 mm. high.

As this species does not appear to have been described, it may be known as *Cordyceps Erotyli*. It is quite unlike *C. curculionum*. The conidial stage may be named *Spicaria Erotyli*.

Cordyceps Erotyli Petch, n.sp.—Clavis numerosis, usque 15 mm. longis, simplicibus vel furcatis; stipitibus 0.15–0.25 mm. diam., externe pruinosis flavis, interne subtranslucidis, rubro-brunneis; capitibus clavatis vel cylindraceis, 1.5–4.5 mm. longis, 0.36–0.75 mm. diam.; peritheciis superficialibus, conoideis vel ampullaceis, 0.4 mm. alt., 0.18 mm. diam., confertis, interdum ab uno latere capitis deficientibus, succineis dein brunneo-rubris, leniter tomentosis; ascis cylindraceis, capitatis, 250 \times 4 μ ; ascosporis linearibus, 0.75 μ diam. On *Erotylus* sp. (Coleoptera), Trinidad.

Spicaria (Isaria) Erotyli Petch, n.sp.—Conidiophoris sparsis vel in clavis albis laxis minutis usque 1.5 mm. alt. aggregatis, simplicibus, usque 130 μ alt., 6 μ diam., phialidibus in verticillis duobus vel tribus

capitulum $25-45\mu$ alt. formentibus; prophialidibus nullis; phialidibus ampullaceis, $6-14 \times 2-2.5\mu$; conidiis cylindratis, utrinque rotundatis, $3-5 \times 1-1.5\mu$. On *Erotylus* sp. (Coleoptera), Trinidad.

109. *CORDYCEPS RAMOSA* Petch, n.sp.

Two collections of this species, made in Trinidad by J. B. Rorer and sent to Thaxter, are now in the Farlow Herbarium, Nos. 6133 and 2626. They were said to be on lepidopterous larvae, but that was queried by Thaxter, and the larvae appear to be coleopterous.

In each specimen, two clavae arise from the larva, which is covered with a thin layer of rufous mycelium. The clavae are up to 7 cm. high, ascending, 1 mm. diameter below, attenuated upwards, longitudinally sulcate, terete, ashy brown, glabrous, and bear up to six lateral branches which arise more or less at right angles to the main stem and curve upwards, sometimes branching again. The main stem and the branches are covered with scattered or crowded, ovate perithecia, 0.4 mm. high, 0.3 mm. diameter, glabrous, with a subacute apex. The perithecia on the branches are now dark brown, subtranslucent, collapsing laterally, while those on the main stem are rufous brown and rigid. The structure of the perithecium is the same in both situations, but only the perithecia on the branches contain asci, those on the main stem being filled with hyphae, an occurrence which is not uncommon in *Cordyceps*. The wall of the perithecium is brown, and composed exteriorly of large polygonal cells, from $9 \times 7\mu$ to $18 \times 15\mu$. The asci are narrow clavate, $130-170 \times 6-7\mu$, and the ascospores $1.5-2\mu$ diameter, with septa $6-7\mu$ apart. Neither free ascospores nor part-spores have been seen, and consequently there is a possibility that the fungus may be an *Ophiocordyceps*. In No. 2626 the perithecia are immature.

Besides its branched character, this species differs from *O. acicularis* in the nature of the main stem, the shape of the perithecia, and the length of the ascus. In general shape it resembles *Cordyceps Henleyae* Massee, but differs from that species in the shape of the perithecia.

Cordyceps ramosa Petch, n.sp.—Clavis usque 7 cm. alt., basi 1 mm. diam., sursum attenuatis, ramis lateralibus usque sex ferentibus, teretibus, cinereo-brunneis, glabris; peritheciis brunneis, superficialibus, ovatis, apice subacutis, glabris, 0.4 mm. alt., 0.3 mm. diam.; ascis angusto-clavatis, $130-170 \times 6-7\mu$; ascosporis $1.5-2\mu$ diam., septis $6-7\mu$ distantibus. On coleopterous larvae, Trinidad.

110. *CORDYCEPS VARIABILIS* Petch, n.sp.

This species occurs on beetle larvae in rotten wood. There are several specimens in the Farlow Herbarium, ex Herb. Thaxter, differing to such a degree that it is difficult to draw up a description which will cover all the forms.

In No. 613 A, on an elaterid larva, Ithaca, N.Y., 25 July 1916, coll. G. H. Duff, the clava is up to 1 cm. high. The stalk is about 0.6 mm. diameter, ochraceous, coarsely pruinose or furfuraceous. The perithecia are fused laterally in groups, 1.8–2.5 mm. long, either terminal or subterminal, or lateral at about half the height of the clava, almost or quite encircling it. The groups of perithecia are slightly pulvinate, strongly rough with the apices of the perithecia. The separate perithecia are 0.4 mm. high, 0.25 mm. diameter, flask-shaped with a conical or subcylindric apex, coarsely ochraceous pruinose almost to the apex, which is now translucent red.

In No. 6144, on a Coleopterous larva in wood, Burbank, Tennessee, 7 August 1886, the perithecia are fused in an apical plate almost surrounding the apex of the clava, which is visible only on one side. The stalk is longitudinally striate, matt, not furfuraceous, and the apices of the perithecia are conical. At the base of the clavae are white tufts of mycelium, which contain hyaline, cylindric conidia, straight or slightly curved, with obtuse ends, $10-12 \times 2-2.5\mu$.

In three numbers, it was suggested that the fungus grew on dipterous larvae or tipulid larvae feeding in rotten wood. The hosts, however, appear to be very small Coleopterous larvae. These numbers are No. 6146, in rotten spruce log, [?] Swamp, York, Maine (no date); No. 6138, in rotten bark of *Tilia*, Kittery Point, Maine, 20 August 1920; and No. 2843, in the same locality and substratum, 1895. In these specimens, the clavae project from the wood, only to a height of about 3 mm. or less. The stalk is about 0.12 mm. diameter, pale yellow, minutely pruinose, and bears, when fertile, a small group of about four perithecia at or near the apex. The perithecia are almost free, fused together, as a rule, only at the base. Most of the clavae in these specimens bear minute acervuli, arising anywhere on the clava as white tufts and becoming subtranslucent, yellow, pulvinate, about 0.4 mm. diameter, and containing cylindric, hyaline conidia, slightly curved, $5-6 \times 2\mu$. It is probable that these acervuli are those of a fungus parasitic on the *Cordyceps*. Evidence has been found that the conidial stage of the *Cordyceps* is a *Hirsutella*, but the specimens are not in a fit condition for description.

Two other specimens in the Farlow Herbarium, ex Herb. University of Wisconsin, are marked respectively "*Cordyceps* in rotten wood, Blue Mounds, Wis., August 7, 1905", and "*Cordyceps*, stroma yellowish, 0.5–1 mm. \times 5–8 mm., Blue Mounds, Wis., August 10, 1906". These are small forms on beetle larvae, resembling those in Nos. 6146, 6138 and 2843. The second is accompanied by a slide which shows perithecia with a cylindrical apex, and cylindric part-spores, $5-7 \times 1.5\mu$. In No. 6132 A, the asci are cylindric, $280-300 \times 9\mu$, and the part-spores cylindric, $7-9 \times 1.5-2\mu$.

This species resembles in some respects *Cordyceps Baumanniana*

P. Henn., from West Africa, which was said to be on a caterpillar. The type specimen appears to be on a beetle larva. Hennings figured the lateral head as strongly verrucose with the projecting apices of the perithecia, but he described it as punctulate, and according to my notes on the type the head resembles that of *C. dipterigena*. From *Ophiocordyceps macularis* Mains, this species differs in the shape of the perithecia and the ascospores.

Cordyceps variabilis Petch, n.sp.—Clavis usque 1 cm. alt.; stipitibus usque 0.6 mm. diam., teretibus, sursum attenuatis, ochraceis, crasse pruinosis vel furfuraceis; peritheciis in lamellis subpulvinatis verrucosis, stipitem fere cingentibus, apicalibus vel lateralibus, usque 2.5 mm. longis, aggregatis; peritheciis ampullaceis, 0.4 mm. alt., 0.25 mm. diam., fere ad apicem ochraceo-pruinosis, apice conico vel cylindrico, translucente, rubro-brunneo; ascis cylindraceis, capitatis, 280–300 × 9μ; articulis ascosporarum cylindraceis, 5–9 × 1.5–2μ. On larvae of Coleoptera, North America.

III. *CORDYCEPS MELOLONTHAE* (Tul.) Sacc.

An account of this species, based on the specimens then available, was given in *Trans. Brit. Mycol. Soc.* xix (1935), 165. Since then, further examples have been examined, and these extend the previous idea of this fungus. All the fertile specimens previously examined were more or less obese forms, similar to that figured by Hard as *Cordyceps herculea*, and some difficulty was experienced in understanding how the long, slender, barren clavae, illustrated by Cist and Fougeroux de Bondaroy, could develop into a *Cordyceps* with a stout stalk and a head about 1 cm. diameter. That difficulty has been removed by specimens from British Guiana, which show that *C. Melolonthae* can assume long, slender forms as well as the more obese forms.

The specimens referred to were collected at Vryheid, British Guiana by Dr D. H. Linder (Linder 971). They were found at the base of a palm tree, in loose elevated soil, 5 cm. of the fungus projecting. The collector's note gives the length of the stalk as 10 cm., with a diameter of 4–6 mm., and the length of the head as 2.5–3 cm., with a diameter 7–7.5 mm. A dried specimen has a stalk, 8 cm. long, 2.5 mm. diameter below, expanding to 3 mm. above, and a head, 2 cm. long, 5 mm. diameter. The head is cylindric, somewhat flattened laterally, abruptly distinct from the stalk, with a V-shaped groove at the base on each side, and sometimes slight indications of a previous longitudinal groove. The colour when fresh was "head apricot yellow to Mikado orange; stalk light buff to almost white".

In *Trans. Brit. Mycol. Soc.* xviii (1933), I referred *Cordyceps Rickii* Lloyd to *C. martialis* Speg. That reference was certainly incorrect, for Lloyd's photograph shows the head of *C. Rickii* abruptly distinct from the stalk, whereas in *C. martialis* the stalk merges gradually into the

head. From the photograph of *C. Rickii*, the fungus is apparently on a melolonthid, and its colour when dry was Mars yellow, according to Lloyd. It would seem most probable that *C. Rickii* is *C. Melolonthae*.

As previously stated, it has been customary in North America during recent years to refer to the "White Grub" *Cordyceps* as *C. herculea* (Schw.) Sacc., on the supposition that it was identical with *Sphaeria herculea* Schw. Lloyd, however, examined the type specimen of *S. herculea*, and found that it was a Gasteromycete, *Cauloglossum transversarium* Fr. That fact was known to Ravenel many years earlier, for when he issued *Cauloglossum transversarium* in *Fung. Car. Exsicc.* v, 79, he added the synonym "*Sphaeria herculea* Schwein!", with the personal sign that he had examined the type specimen.

Coker & Couch, in *The Gasteromycetes of the Eastern United States and Canada* (1928), p. 56, remark that "The specimen labelled *Cauloglossum transversarium* in the Philadelphia Herbarium (Ravenel, *Fung. Car. Exs.* 79) is not this species but *Sphaeria herculea* Schw., according to Ravenel." That, however, is not what Ravenel meant. What he intended to convey was that it was *Cauloglossum transversarium* and also *Sphaeria herculea*, the latter name being a synonym of the first. The corresponding specimen under the same number and label in Herb. British Museum is certainly *C. transversarium*, and there are seven other gatherings by Ravenel of that species, correctly named, in the same herbarium.

Mains (*Proc. Amer. Phil. Soc.* LXXIV (1934), 265), perhaps misled by Coker & Couch's misunderstanding, has suggested that the name, *Cordyceps herculea*, should be retained for the "White Grub" fungus, on the ground of general usage. It is difficult to see how that can be done, in view of the valid prior name, *C. Melolonthae* (1865). *C. herculea*, as applied to the "White Grub" fungus, apparently dates only from Ellis and Everhart, *North American Pyrenomycetes* (1892).

It may be noted that determinations of the "White Grub" fungus as *C. Ravenelii* (Riley, *American Entomologist*, III, 137; Masee, *Ann. Bot.* IX, 34) are certainly incorrect, and were based on immature examples of *C. Melolonthae*.

112. *OPHIOCORDYCEPS CALOCEROIDES* (B. & C.) Petch.

In redescribing this species in *Trans. Brit. Mycol. Soc.* XVIII (1933), 63, it was suggested that the reason why the type specimen was collected without its host and was recorded as occurring on the ground was no doubt that the specimen had fallen to the ground and the host had disintegrated.

Further specimens have indicated that the foregoing explanation is probably erroneous. The fungus may occur on a spider which makes a cylindrical hole in the ground and lines it with silk. When attacked

by the fungus, the spider dies at the bottom of its tube, and the fungus may grow up from it along the silk sides of the tube, not in stout strands, but as a delicate mycelium, the clavae arising only near the upper edge of the tube. The fungus would thus appear to be growing on the ground, and unless it was recognized as a *Cordyceps* and the tube carefully dug up, it would be collected without any indication of attachment to a subterranean host.

These recent examples also show that *Ophiocordyceps caloceroides* does not always take the long linear form of the type specimen. One, "on a ground spider, Grand Etang, Grenada, B.W.I., 1912-13" (Thaxter), Farlow Herb. 6122, bears two clavae, with a stalk about 8 mm. high, 0.25-0.4 mm. diameter, and an irregularly ovoid head, rounded above, 1.5-2 mm. high, 1 mm. diameter, abruptly distinct from the stalk. Another, immature specimen from Pehata, Liberia, October 1926 (Linder 1093), also on a ground spider, has at the top of the tube eleven clavae, which are narrow clavate, up to 15 mm. high, 0.25 mm. diameter at the base, expanding uniformly upwards to 0.75 mm. diameter.

113. *CORDYCEPS CYLINDRICA* Petch, n.sp.

This species, which occurs on a trap-door spider, was collected by Dr D. H. Linder in St Anne's Valley, Trinidad, 22 August 1923 (Linder 111). When fresh the head was pale yellow and the stalk white, but the dried specimen is black. The spider, covered with white mycelium, is fixed at the base of the tube, and the single *Cordyceps* clava arises from the spider and emerges at the side of the trap-door.

The stalk is about 2.5 cm. long, 2 mm. diameter, irregularly twisted. The head is cylindric, 1.4 cm. long, abruptly broader than the stalk, 5 mm. diameter at the base, rapidly diminishing to 3 mm. diameter, and thence uniformly cylindric to the obtuse apex. The head is minutely rough with the ostiola, which barely project, and pruinose between them. The perithecia are deeply immersed, flask-shaped with a long neck, 1.2 mm. high, 250 μ diameter. The head is red-brown in section, soft and subgelatinous internally, and, judging from the grains of sand adhering to it, it is viscid when fresh. Unfortunately, no asci were found, the specimen being apparently effete. A few spores, cylindric or narrow-oval with rounded ends, 2.5-3.5 \times 1 μ were observed, but it is doubtful whether these were part-spores.

It seems probable that this species is related to *Isaria atypicola* Yasuda, described from specimens on a spider in Japan, which is the same shape, and is said to be light purple above, velvety, with cylindrical conidia.

Cordyceps cylindrica Petch, n.sp.—Stipite ca. 2.5 cm. alt., 2 mm. diam., irregulariter torto, albo, sicco nigro; capite cylindraco,

1.4 cm. alt., a stipite distincto, basi 5 mm. diam., ad 3 mm. diam. mox attenuato, dein ad apicem cylindraceo, pallide flavo, sicco nigro, pruinoso, ostiolis minute scabro, interne rubro-brunneo, subgelatinoso; peritheciis immersis, ampullaceis, collo longo, usque 1.2 mm. alt., 250μ diam. Ascis et sporis non visis. On a trap-door spider, Trinidad.

114. *CORDYCEPS ELONGATA* Petch, n.sp.

In the Farlow Herbarium there are four specimens of this species, ex Herb. Thaxter, on the larva and pupa of *Apatela americana*, No. 6134, Burbank, Tennessee, September 1887; No. 6135, Cranberry, North Carolina, 1887; No. 6137, York, Maine, August 1893; and No. 6277, York, Maine, and Cullowhee, N.C. No. 6137 is the type.

The clavae are up to 11 cm. high, from one to three arising from each host. The stalk is flexuose, longitudinally sulcate and twisted, pale brown, nearly glabrous, about 1 mm. diameter, almost equal, or expanding upwards to 2 mm. diameter. The head is from 1 to 4 cm. long, slightly thicker than the stalk, 1.1–2.2 mm. diameter, cylindric, terete, equal, or in the thicker forms slightly attenuated upwards, obtuse, or rarely with a minute apiculus, pale yellow, rough with red-brown ostiola. Sometimes the perithecial area is lacking in a narrow strip down one side of the head. The perithecia are immersed, the tissue between their apices forming a thin, continuous sheet on the dried specimens; they are scattered or crowded, ovato-conoid, 0.5 mm. high, 0.3 mm. diameter, apex subacute, wall yellow by transmitted light. The asci are 220μ long, 8μ diameter, and the ascospores 2μ diameter, with septa 4–12 μ apart. Part-spores were not seen, but the ascospores are cylindric, and the fungus is probably *Cordyceps*, not *Ophiocordyceps*.

This species resembles a thin, elongated *Cordyceps militaris*. It has, however, a different stalk, and its asci and ascospores are broader than in that species. The colours described above are those of the dried specimen. From similarity of structure of the stalk, it seems probable that its conidial stage is *Hirsutella gigantea*, described in the next note.

Cordyceps elongata Petch, n.sp.—Clavis paucis, usque 11 cm. alt.; stipitibus flexuosis, longitudinaliter striatis, tortis, pallide brunneis, fere glabris, circa 1 mm. diam., aequalibus vel sursum incrassatis ad 2 mm. diam.; capitibus 1–4 cm. alt., stipite vix crassioribus, 1.1–2.2 mm. diam., cylindraccis, teretibus, pallide flavis, ostiolis rubro-brunneis asperis, apice obtuso vel rarius breviter apiculato; peritheciis immersis, sparsis vel congestis, ovato-conoideis, 0.4 mm. alt., 0.3 mm. diam., apice subacuto, pariete flavo; ascis cylindraccis, $220 \times 8\mu$; ascosporis 2μ diam., septatis, septis 4–12 μ distantibus. On pupae and larvae of *Apatela americana* (Lepidoptera), North America.

115. *HIRSUTELLA GIGANTEA* Petch, n.sp.

The type specimen of this species, Farlow Herb. 6126, was collected by Thaxter "Above Diana's Bath, Intervale, New Hampshire, July 1901." It is on a Lepidopterous pupa attached to a piece of wood, and bears three clavæ, all of which have been broken. Two of them have regrown, but the apices are lacking in all three. In their present condition, the height is about 4 cm. They are about 0.6 mm. diameter, brown below, longitudinally sulcate, glabrous, ashy and minutely setose above. The upper part of the clava is covered by a palisade layer of clavate or cylindric cells, rounded above, $15-18 \times 6-7\mu$, among which are *Hirsutella* phialides up to 40μ high, with a flask-shaped base, $16-20 \times 8-9\mu$, and a long, stout sterigma, 1μ diameter. The spore cluster is lemon-shaped, $10 \times 6\mu$, becoming globose, 10μ diameter, and the separate conidia are broadly cymbiform with obtuse tips, $9-10 \times 3-4\mu$.

In another specimen, Farlow Herb. 6199, on a cocoon of *Apatela*, R. Thaxter, York, Maine, about fourteen clavæ, now all pale brown, arise from the cocoon. These are straight below, flexuose above, with scattered lateral branches. The specimen has been pressed, and now covers an area 8.5 cm. high and 7 cm. wide, while the clavæ are so entangled that it is not possible to ascertain their exact height, but some of them certainly exceed 10 cm., though their diameter at the base is only 0.5 mm.

In contrast to the foregoing specimen is one (Farlow Herb. 648), collected by Thaxter at New Hartford, Conn., 1888-9, in which the clavæ are only about 2 cm. long. These are on a small pupa, about 1 cm. long.

The bases of the branched stalks in the specimen of *Stilbella ramosa* (Peck), Farlow Herb. 6142, described later in this series, exactly resemble the stalks of *Hirsutella gigantea*.

There would seem to be some possibility that this species may be *Isaria nigripes* Schw., *North American Fungi*, No. 3068. Schweinitz stated that his species grew on buried chrysalides, and was simple, half an inch high, with a somewhat slender, black, glabrous stalk up to half the height, passing into a simple, obtuse, ashy white, pulverulent head, often falcate, the conidia rather compact. But Schweinitz's type does not appear to have been re-examined, and the fungus has not been reported since.

Hirsutella gigantea Petch, n.sp.—Clavis 2-10 cm. et ultra alt., 0.5-0.6 mm. diam., simplicibus, vel supra ramosis, teretibus, saepe longitudinaliter sulcatis, infra brunneis, pruinosis dein glabris, supra cinereis, setosis; phialidibus ad 40μ alt., basi ampullacea, $16-20 \times 8-9\mu$, sterigmate longo, crasso, 1μ diam.; conidiis late cymbiformibus,

obtusis, $9-10 \times 3-4\mu$. On pupae of Lepidoptera (*Apatela americana*, etc.), North America.

116. *CALONECTRIA HIRSUTELLAE* Petch, n.sp.

In *Trans. Brit. Mycol. Soc.* xvi (1932), 226, I described *Calonectria pruinosa*, the ascigerous stage of *Hirsutella versicolor*, a common *Hirsutella* on leaf-hoppers in Ceylon. The corresponding *Hirsutella* on leaf-hoppers in the Western Hemisphere is *H. floccosa* Speare, and in the Farlow Herbarium there is a specimen, No. 6153, collected by Thaxter at Cranberry, North Carolina, August 1887, bearing the ascigerous stage, which again is a *Calonectria*.

The insect is covered with a compact, white or cream-coloured film of mycelium which sometimes spreads out in a radiating mat over the leaf. This mycelium bears *Hirsutella* conidiophores. The perithecia are semi-immersed, globose, 0.25 mm. diameter, with a short, cylindric ostium, 0.05 mm. high, apex obtuse. They are now dark amber, glabrous above. The asci vary from clavate, $80 \times 12\mu$, spores biseriata above, uniseriate below, to linear clavate, $130 \times 6\mu$, spores uniseriate. The apex of the ascus is thickened, but not with a long, truncate, solid apex as in *Calonectria pruinosa*. The asci are eight-spored, and are accompanied by ligulate, diffuent paraphyses. The ascospores are fusoid, straight or slightly curved, ends obtuse, seven-septate, $22-27 \times 5-6\mu$.

This species differs from *C. pruinosa* in the shape of the perithecia, the apex of the ascus, and the shape of the ascospores. It may be known as *C. Hirsutellae*.

Calonectria Hirsutellae Petch, n.sp.—Mycelio albo, insectum obducente. Peritheciis semi-immersis, globosis, 0.25 mm. diam., collo brevi cylindrico, 0.05 mm. alt., apice obtuso, succineis (sicco), supra glabris; ascis octosporis, aut clavatis, sporis supra biseriatis, infra uniseriatis, $80 \times 12\mu$, aut lineari-clavatis, sporis uniseriatis, $130 \times 6\mu$, apice incrassato; paraphysibus ligulatis, diffluentibus; ascosporis hyalinis, fusoidis, rectis vel leniter curvatis, utrinque obtusis, septem-septatis, $22-27 \times 5-6\mu$. On a leaf-hopper, Cranberry, North Carolina.

117. *TORRUBIELLA PAXILLATA* Petch, n.sp.

In the Farlow Herbarium, ex Herb. Thaxter, there are three gatherings of a fungus "on aphid lion", No. 6148, Cranberry, North Carolina, 1886; No. 6150, Burbank, Tennessee, 1898; and No. 6151, Cranberry, North Carolina, August 1887. An aphid lion, I am informed, is the larva of a lace-wing fly (Chrysopid, Neuroptera). The fungus is a *Torrubiella*.

The insect, or rather the debris over the insect, is covered by a

delicate, white film of mycelium, tomentose with short, erect conidiophores. The perithecia of the *Torrubiella* are flask-shaped, 0.6 mm. high, 0.25 mm. diameter, contracted towards the base into a peg, partly immersed in the loose mycelium, now amber-coloured, minutely rugose, naked above, or with scattered, white tufts of mycelium. The asci are cylindric, capitate, $280-330 \times 5\mu$, and the ascospores linear, 1μ diameter, almost as long as the ascus, with septa $7-9\mu$ apart. Free part-spores were not found. No. 6148 has the more nearly mature perithecia, the other two being immature.

The conidiophores are erect, usually simple, rarely branched, 3μ diameter, hyaline, septate, with the terminal segment sometimes oval and inflated to 4μ . In some the segments of the conidiophore vary in breadth, segments 1.5μ diameter alternating with segments 3μ diameter. Nearly all the upper segments are covered with lateral, obtuse, cylindric sterigmata up to 1μ high, but these are less numerous on the thinner segments and may be lacking there. The conidia are oval or pyriform, hyaline, $2-4 \times 1.5\mu$. This conidial form is similar to *Gonatorrhodiella coccorum* Petch, from which it differs in the shape of the conidiophore, and the size and shape of the conidia.

Torrubiella paxillata Petch, n.sp.—Mycelio insectum bysso albo obducente; peritheciis ampullaceis, 0.6 mm. alt., 0.25 mm. diam., infra contractis, paxillatis, nonnihil immersis, succineis, minute rugosis, supra glabris; ascis cylindraceis, capitatis, $280-330 \times 5\mu$; ascosporis linearibus, 1μ diam., septatis, septis $7-9\mu$ distantibus. On the larva of a lace-wing fly (Chrysopid, Neuroptera), North America.

118. *TORRUBIELLA GONYLEPTICIDA* (Möller)

In a collection of entomogenous fungi made in the West Indies by Dr C. B. Williams, there was one on a fairly large spider, Trinidad, December 1916. The spider was fastened to a fragment of bark by white mycelium, which sparsely covered the body and legs, and here and there formed a white or cream-coloured felted crust, somewhat nodular, especially on the abdomen. Traces of a *Spicaria*, with flask-shaped, long-necked phialides, and oval conidia, $2-3 \times 1.5-1.75\mu$, were found, but the specimen was not adequate for description.

Recently, it has been possible to examine further specimens. In one, collected by Thaxter, Maravel Valley, Trinidad, 1912-13 (Farlow Herb. 2569), the body and legs of the spider are covered by a very delicate, white film of mycelium. Another, on a large spider, collected by J. B. Rorer, Trinidad (Farlow Herb. 2586), and one on *Mygale*, collected by Thaxter, Sangre Grand, Trinidad (Farlow Herb. 2570), bear conidiophores, but are in poor condition. The best example available was collected by Thaxter, on a large spider, Maravel Valley, Trinidad, 13 February 1913 (Farlow Herb. 2559).

The fungus covers the body and legs of the spider with a delicate white film of mycelium, which then produces a dense pile of erect conidiophores over the body, and a looser growth of the same conidiophores along the legs. The covering on the body appears granular owing to the *Spicaria* heads. It is at first white, becoming cream-coloured, on the body, the looser growth on the legs appearing greyish, with cream-coloured clusters of conidia. Here and there it produces erect, cylindric or clavate, loose synnemata, up to 3 mm. high. The conidiophores are up to 360μ high, with a stalk 3μ diameter, and have two or three whorls of prophialides, or of short branches, in the uppermost $40-60\mu$. The prophialides are ovate, or cylindric constricted towards the middle, $5-7 \times 2.5-3\mu$, and the phialides flask-shaped with a long neck, up to 9μ long, 2.5μ diameter below. The conidia are narrow-oval, sometimes with one end subacute, $2-5 \times 1.5-3\mu$, or subglobose, $2-3\mu$ diameter, catenulate. This species may be known as *S. longipes*.

Spicaria longipes Petch, n.sp.—Conidiophoris confertis, albis dein flavescentibus, usque 360μ alt., stipite 3μ diam., prophialides vel ramos breves in duobus vel tribus verticillis supra ferentibus; prophialidibus ovatis, vel cylindraceis medio constrictis, $5-7 \times 2.5-3\mu$; phialidibus ampullaceis, $9 \times 2.5\mu$, collo longo; conidiis anguste ovalibus, hyalinis, continuis, interdum uno fine subacutis, $2-5 \times 1.5-3\mu$, vel subglobosis, $2-3\mu$ diam., catenulatis. On spiders, Trinidad.

Another specimen was collected by Thaxter in Maravel Valley, Trinidad, 1912-13 (Farlow Herb. 2563), and was marked by him, "on large spider under logs, apparently perfect form of the large, white spider *Isaria*." The body of the spider in the dry state measures about 7×5 mm., and the spread of its legs is about 25 mm. The spider is sparsely covered with white mycelium, and in a powdery patch on the upper side of the body are densely clustered conidiophores which agree with the foregoing description.

Scattered over the body and legs of this last-mentioned specimen are the perithecia of a *Torrubiella*. These are superficial, ovato-conoid, obtuse, 0.4 mm. high, 0.27 mm. diameter, minutely rugose, with a few hyphae at the base, and sometimes white pruinose. Their colour is now dark amber. The perithecial wall bears large, oval or subglobose, hyaline cells, up to 25μ diameter, sometimes in short chains; these are external to the wall proper, which is transversely hyphal, as usual in *Torrubiella*.

This appears to be referable to *T. gonylepticida* (Möller). Möller's fungus was on a small spider, found among fragments of bark on the ground. He stated that the perithecia were $300-400\mu$ long and had a comparatively thick plectenchymatous wall, the outer cells of which were fairly isodiametric, and projected so that the wall appeared uneven and obtusely nodulose. On the other hand, Möller also stated

that there was not the smallest trace of mycelium on the body of the spider. This latter statement, however, must be accepted with caution, and even if true it is scarcely possible that the absence of superficial mycelium would be a constant character. On some of the specimens recorded here the mycelial covering is so slight that it might be overlooked if the specimen were preserved in alcohol. The structure of the perithecial wall would appear to be a more decisive character than the presence or absence of mycelium on the host.

If this identification is correct, *Spicaria longipes* is the conidial stage of *Torrubiella gonylepticida*.

119. *STILBELLA KERVILLEI* (Quélet) Lingelsh.

Stilbella Kervillei was described by Quélet (as *Stilbum*) from specimens on the fly, *Leria (Blepharoptera) caesia*, found in limestone caves near Rouen and Elbeuf, and it has since been found in similar caves at Maastricht on *Blepharoptera* spp. and *Scoliocentra* spp. It was recorded for England by Mr F. A. Mason in *Journal of Botany*, August 1931, pp. 205-7, having been discovered by Mr Leslie Armstrong on *Blepharoptera serrata* in the Creswell Caves, Derbyshire.

Quélet described his species as having a white, villous stalk, often branched or proliferous, a yellow, globose head, and brown mycelium on the insect. The British specimens recorded by Mr Mason had simple stalks, but otherwise agreed with Quélet's description. The insect was partly covered by a thin, discontinuous, rufous brown crust of mycelium, in which occurred dark brown, lemon-shaped bodies, resembling the spore clusters of a *Hirsutella*, but it was not possible to separate these into distinct conidia, nor to detect *Hirsutella* phialides.

With the flies which bore the *Stilbella* were others bearing long, flexuose, hair-like, brown filaments, from 2 to 6 cm. long and 0.1 mm. diameter, which were sterile and resembled the sterile clavae which occur in some species of *Hirsutella*.

In March 1934, Mr Armstrong collected more fungi on flies in Pinhole Cave, Creswell. These included branched examples of *Stilbella Kervillei*, another example of the long, thin clavae, and a single specimen which proved to be a typical *Hirsutella*.

The last-named specimen bears four *Hirsutella* clavae. These are up to 8 mm. long, 0.2 mm. diameter, simple, terete, tapering to an acute apex, rigid, ashy, slightly pruinose, fuscous internally. The phialides are up to 27μ high, with a conoid base, $4-6 \times 3-4\mu$. The spore cluster is lemon-shaped, about $10 \times 6\mu$, and the individual conidia are greenish hyaline, cymbiform, ends acute, $6-10 \times 1.5-2\mu$. The insect is partly covered by a thin, rufous brown crust of mycelium, and this contains *Hirsutella* phialides, and dark brown spore clusters, identical with those found in the brown mycelium which accompanies *Stilbella Kervillei*. I propose to call this species *Hirsutella dipterigena*.

The conidial cluster of a *Hirsutella* consists of a small number of conidia, usually parallel to one another, embedded in mucilage. In some species the adherence of the conidia is only slight, and they soon separate and may be found adhering to the lower part of the sterigma or to the mycelium. In others the cohesion is greater, but the conidia separate in water. In the present species the conidia adhere strongly to one another, and the spore cluster persists as a whole and turns dark brown when old, as in *Synnematium Jonesii*.

There is an example of the long, hair-like, sterile form of this *Hirsutella* in Herb. British Museum, on *Blepharoptera serrata*, found in a "stalactite cave", Yealhampton, Devon.

Hirsutella dipterigena Petch, n.sp.—Mycelio rufo-brunneo, insectum strato tenui interrupto obducente; clavis usque 8 mm. alt., 0.2 mm. diam., simplicibus, teretibus, rigidis, cinereis, leniter pruinosis, interne fuscis; phialidibus usque 27μ alt., basi conoideo, $4-6 \times 3-4\mu$; conidiis viridi-hyalinis, cymbiformibus, acutis, $6-10 \times 1.5-2\mu$, in pseudosporis limoniformibus, circa $10 \times 6\mu$, muco conglomeratis. On *Blepharoptera serrata*, Pinhole Cave, Derbyshire.

The branched examples of *Stilbella Kervillei* had a main stem up to 16 mm. long, sometimes with an acute tip, sometimes terminating in a *Stilbella* head. The lateral branches were short and simple, usually furnished with a globose head, but the branching was irregular, the laterals being sometimes scattered along the main stem and perpendicular to it, sometimes clustered, two or three together, at varying angles. On mounting the whole of one of these branched clavae, *Hirsutella* phialides were found on the main stem, showing that it was a *Hirsutella* clava. Thus, *Stilbella Kervillei* is not parasitic on the fly, but on an entomogenous fungus, *Hirsutella dipterigena*. The brown mycelium on the fly from which the simple *Stilbella* arises is that of the *Hirsutella*.

In *Bull. Soc. Nat. Sci. Buffalo*, 1 (1873), Peck described *Stilbum ramosum* as "Head subglobose, whitish or pale yellow; stem thick, smooth, branched, white above, pallid or brownish below, sometimes creeping and sending up branches at intervals; spores minute, oblong. Dead larvae of insects buried in rotten wood." In current nomenclature this would be *Stilbella ramosa*.

I have not seen the type of *S. ramosa*, but in the Farlow Herbarium there are several specimens which appear to be referable to Peck's species. No. 6142, Burbank, Tennessee, 1896, is on a larva, apparently Lepidopterous, which bears seven scattered clavae, 7.5–14 mm. high, two of which are branched. The stalk is stout, 0.2–0.5 mm. diameter, almost equal, now pale brownish, dark brown at the base, longitudinally furrowed, minutely pruinose, becoming glabrous. The head is globose or ovoid, up to 1.25 mm. diameter, yellow, waxy. The conidiophores in the head are linear, branched, and the conidia oval, oblong-oval, or pyriform, $2-3.5 \times 1-1.5\mu$.

Another specimen, Farlow Herb. 6135, Cranberry, N.C., 1887, also on a larva, is a new species of *Cordyceps*, *C. elongata* Petch, one clava of which has examples of *Stilbella ramosa* growing on the stalk. These are 1–1.5 mm. high, with a white, terete stalk, 0.36 mm. diameter at the base, 0.12 mm. diameter above, minutely tomentose, and a yellow, ovoid head, about 0.36 mm. high, 0.3 mm. diameter. The conidia are the same as in the previous specimen. Thus, in this instance, *S. ramosa* is a simple *Stilbella*, and it is parasitic on a *Cordyceps*.

A third specimen, Farlow Herb. 646, Woodmont, Conn., September 1888, is on a small Lepidopterous pupa, which bears about ten *Stilbella* synnemata, up to 5 mm. high, with stalks 0.15 mm. diameter, white, minutely tomentose, and dark orange, subglobose heads, 0.4 mm. diameter. The conidia are oval or subpyriform, $1.5\text{--}2.5 \times 1\text{--}1.5\mu$, or globose, 1.5μ diameter. The pupa also bears the bases of dark brown, linear clavae, two of which have regrown, and these are the clavae of *Hirsutella gigantea*. Thus, here, *Stilbella ramosa* is parasitic on a *Hirsutella*.

Comparison of No. 6142 with No. 6135 and other examples of the same *Cordyceps* shows that the apparent stalks of the *Stilbella* in No. 6142 are really the broken stalks of *Cordyceps elongata* or *Hirsutella gigantea*, only the apices and the lateral branches being the *Stilbella*, which is actually a simple synnema.

Thus, *Stilbella ramosa* (1873) is similar to *S. Kervillei* (1884) in being parasitic on a *Hirsutella* or on a *Cordyceps* of which that *Hirsutella* is the conidial stage. The differences between the American and the European *Stilbella* are slight, and I am of opinion that they are the same species. *S. Arndtii* Lingelsh., found on *Blepharoptera serrata* in caves in Silesia, is doubtless the same species.

120. *ISARIA SPHECOPHILA* Ditm.

Isaria sphecophila was described and figured by Ditmar in Sturm's *Dtsch. Flora*, Abt. III, Bd. I, p. 115, pl. 57. It grew on a hornet, which lay in a cavity in a dried stem. Ditmar's figure shows fifteen linear clavae, up to 9 cm. long, springing from the insect, most of them with a knot, or irregularly thickened region, at about one-third their height from the base. Ditmar described the clavae as "medio nodosis", but it would appear that that feature was accidental, probably caused by contact of the clavae with the edge of the cavity. He also described the upper part of the clava as pilose, and figured it minutely setulose in his drawing of part of the clava enlarged. This last character has caused some difficulty or doubt in the identification of Ditmar's species. As pointed out by Speare (*Mycologia*, XII, 63), Ditmar's figure of a setulose clava suggests that his fungus was a *Hirsutella*. On the other hand, the available specimens in herbaria which would otherwise be referred to *Isaria sphecocephala* are *Hymenostilbe*.

In September 1933 I collected an "*Isaria*" on an ichneumon at Chopwell Wood, Northumberland. The clavæ were small, the largest being only 1.75 mm. high, subulate, 0.4 mm. diameter at the base, 0.2 mm. diameter above, apex subacute, cream-coloured. Under a hand lens, the clava was very minutely setulose. Nevertheless, the fungus was a *Hymenostilbe*, the setulose appearance being due to projecting phialides and their attached conidia. The phialides were flask-shaped, $12 \times 4\mu$, and the conidia clavate, the upper end rounded, the lower truncate, hyaline, smooth, $6-10 \times 4-5\mu$.

In a specimen, Rehm, Ascomyceten, No. 1287, "*Cordyceps spheco-phila* (Kl.) B. & C.", there was a conidial clava. This was not setulose under a lens. It bore a palisade layer of phialides, $16-24\mu$ high, 4μ diameter, clavate, with a truncate apex. The conidia were clavate, $7-9 \times 3.5-4\mu$. This conidial stage is *Hymenostilbe*.

In a specimen from Porto Rico in the Farlow Herbarium, a similar conidial clava had phialides flask-shaped, or narrow-oval, attenuated at the apex, $13-18 \times 4-5\mu$, and conidia oval or narrow-oval, obtuse, $9-12 \times 4\mu$.

It may be concluded that *Isaria spheco-phila* Ditm. is a *Hymenostilbe*, and that it is the conidial stage of *Cordyceps sphecocephala* (Kl.) Cooke. It will stand as *Hymenostilbe spheco-phila* (Ditm.).

121. *HYMENOSTILBE AMPULLIFERA* Petch, n.sp.

This species is included in the Farlow Herbarium under two numbers, No. 727, "on gnats, coll. R. Thaxter, Cranberry, N.C., August 1887", and No. 6254, "on gnats, coll. R. Thaxter, Cranberry, N.C., 1887", probably parts of the same gathering. No. 727 bears the identification of the insect as a tipulid, *Dicranomyia pubipennis* O.S.

The insects bear small, white, irregularly cylindric or conical clavæ from the joints of the legs, the body, and the tip of the abdomen, as well as minute, white tufts scattered along the legs. The largest clava is cylindric, 3 mm. long, appearing 0.25 mm. diameter because of a loose covering of conidia, but about 80μ diameter without the conidia, and tapering upwards to about 40μ . The central core is composed of agglutinated parallel hyphae, and bears a palisade layer of flask-shaped phialides, $11-15 \times 4-5\mu$, and barren, cylindric or clavate cells, rounded above, not verrucose, $10 \times 3\mu$. The conidia are hyaline, narrow oval, $7-11 \times 2-2.5\mu$, compressed and appearing cylindric, 1.5μ thick, when viewed laterally. This species differs from *Hymenostilbe muscaria* in its phialides and conidia.

Hymenostilbe ampullifera Petch, n.sp.—Clavis minutis, albis, pruinosis, usque 3 mm. alt., 80μ diam., supra attenuatis; phialidibus ampullaceis, $11-15 \times 4-5\mu$, cellulis sterilibus, cylindræis vel clavatis, apice rotundatis, intermixtis; conidiis anguste ovalibus, hyalinis,

7-11 × 2-2.5 μ , a latere compressis et cylindraceis, 1.5 μ crass. On *Dicranomyia pubipennis* O.S. (Tipulidae), North America.

122. *HYMENOSTILBE FRAGILIS* Petch, n.sp.

This species, which occurs on Orthoptera, was collected by Thaxter in Verdant Vale, Trinidad, October 1912, "on an orthopterous larva" (Farlow Herb. 2581). The insect is covered with a yellowish loose film of mycelium, from which arise large numbers of white stilboid processes, erect, usually flexuose, up to 1.3 mm. high, 0.1 mm. diameter at the base, almost equal, or attenuated upwards to 0.05 mm. diameter, expanding into a clavate or ovoid head, up to 0.18 mm. diameter. Both the head and the stalk bear a palisade layer of cylindric phialides, 6-9 μ high, 3 μ diameter, with an obtuse, rounded apex, rough with spherical globules up to 1 μ diameter. The conidia are fusoid or narrow-oval, ends obtuse, 6-10 × 2-3 μ . The outer layer of the clava readily disintegrates under pressure, or when old, leaving a thin, glabrous, horny, linear clava, sometimes pruinose with the remains of the conidial layer. I have examined another specimen of this species on a locustid from British Guiana, collected by E. B. Martyn. I propose to name it *Hymenostilbe fragilis*.

A specimen collected by F. W. Urich, "on a cricket", Concord Estate, Sangre Grande, Trinidad, 4 January 1917 (Farlow Herb. 6162) bears two perithecial clavae and large numbers of conidial clavae. The latter are 0.6-0.8 mm. high, 0.02 mm. diameter below, gradually becoming clavate in the upper half and 0.06 mm. diameter above. Disintegration of the outer layer has set in, and the stalk is yellowish, translucent and horny, but the head is still entire and white. The phialides agree with those of the previous specimens, and the conidia are fusoid, sometimes oblique at the base, with a subacute apex, 8-10 × 2 μ . The conidia are borne on a minute, central sterigma, not on the globular processes. The perithecial clavae in this specimen are *Cordyceps Uleana* P. Henn. Consequently, *Hymenostilbe fragilis* is the conidial stage of *Cordyceps Uleana*.

Another specimen of *C. Uleana*, which bears both perithecial and conidial clavae, is included in the cover of "*Cordyceps Sphingum*" in Herb. Kew., "Traill 23, on Orthoptera, R. Jurua, 7. 11. 74."

Hymenostilbe fragilis Petch, n.sp.—Clavis usque 1 cm. alt., 0.5 mm. diam., rectis vel curvatis, aequalibus vel sursum attenuatis, in capitulo albo clavato vel ovali desinentibus; phialidibus cylindraceis vel ovatis, 6-13 × 3-4 μ , apice rotundatis, supra glandulosis; conidiis fusoides vel clavatis vel ovalibus, obtusis vel uno fine subacutis, 5-12 × 2-3 μ . On Orthoptera, Trinidad, British Guiana, Brazil.

123. *HIRSUTELLA SUBULATA* Petch

The only previously known specimen of this species was that on a caterpillar found at Milton, Northamptonshire, England, and recorded by Berkeley and Broome as *Isaria floccosa* Fr. in *Ann. Mag. Nat. Hist.*, Ser. 5, VII (1881), 130.

In the Farlow Herbarium, No. 4, there is a specimen on the larva of *Aegeria pyri* Harris in apple bark, coll. J. L. Zabriskie, Flatbush, L.I., N.Y., 23 September 1890. The larva bears three clavae, with the bases of four or more others. The clavae are now ashy, becoming pale brown, linear, terete, up to 1.5 cm. high, 0.3 mm. diameter, tapering upwards, minutely pruinose. The conidia measure $4-6 \times 1.5-2\mu$, and the spore cluster is oval, up to $8 \times 5\mu$.

Another specimen (Farlow Herb. 6241), on the larva of the "Codling Moth" under a piece of bark, was collected by Riley at Washington, D.C. The clavae are now broken, but there were apparently about a dozen, up to 7 mm. high, 0.4 mm. diameter at the base, clustered, tapering upwards to an obtuse tip, curved above, white pruinose, internally yellowish and subtranslucent. The spore cluster is oval, $6 \times 3\mu$, and the conidia cymbiform, $4 \times 1.5\mu$.

124. *ISARIA BARBERI* Giard

In *C.R. Soc. Biol.*, Paris, séance du 22 Déc. 1894, p. 823, Giard gave the name *Isaria Barberi* to a fungus on the larva of the Moth Borer (*Diatraea saccharalis* Fabr.), which had been sent to him from the West Indies by Dr C. A. Barber. The yellowish clavae arose from all parts of the larva, and were oriented in the direction of its length, probably because the fungus attacks the insect in its burrow and the clavae grow towards the entrance. The best developed clavae were like a pollarded tree in shape, with a short, basal stem, 5-15 mm. long, 1 mm. diameter, which bore at the apex two to four branches, 3-5 cm. long, 0.6 mm. diameter. The conidiophores were scattered along the clavae, and were cylindrico-conical, $4-5\mu$ high, with one, sometimes two, slender sterigmata, $3-4\mu$ high, terminated by an ovoid spore, 1.5μ long.

In *Ann. Bot.* IX (1895), 18, Massee described a *Cordyceps* on the same insect from Barbados under the name *C. Barberi* Giard, stating that the specimens had been sent to Kew by Mr John R. Bovell during the autumn of 1894, and had been labelled *C. Bovellii*, but the name *C. Barberi* had been adopted as having priority, on the assumption that *Isaria Barberi* Giard was the conidial form of the ascigerous condition described.

In Herb. Kew., in the cover labelled *Cordyceps Barberi*, there are three packets. One of these is marked "*Cordyceps Bovellii* Mass., on

larvae of sugar-cane borer, in sugar cane, Barbados, November 1894", and contains several larvae and clavae, the latter being linear, sometimes with short lateral branches, up to 15 mm. long, 0.25 mm. diameter, straight or variously curved, white or ashy, becoming brown, mostly detached. A second packet is marked "*Cordyceps* sterile, parasitic on insects injurious to sugar cane, Barbados (Barber)", and contains a single larva, with clavae up to 8 mm. long, 0.3 mm. broad, somewhat flattened, dark brown, smooth, matt, with short, thorn-like lateral branches. The third packet is marked "*Cordyceps Barberi* Giard, on larva of borer in sugar cane, Trinidad, Hart, October 1900", and contains apparently a pupa, which bears several short, linear or subulate clavae, white or cinereous, up to 3 mm. long.

Massee gave a figure of *C. Barberi*, but there is nothing now among the specimens which answers to the description and figure, and Dr F. J. Seaver has kindly informed me that there is no specimen of *C. Barberi* in the Massee collection in New York. The specimens in Herb. Kew. are a *Hirsutella*. The base of the phialide is ovate, conoid, or cylindric, $5-7 \times 3-4\mu$, attenuated into a slender sterigma, $7-9\mu$ long. The spore cluster is subglobose or oval, $4-7 \times 3.5-4.5\mu$, and the conidia are narrow-oval or fusoid, $4.5-7 \times 2\mu$, one end acute. On the older parts of the clava, many of the phialides are larger and irregular, with a cylindric, septate base, sometimes branched, probably due to growth after the abscission of the conidia.

Giard's description of the conidiophore is that of a *Hirsutella*. The clavae described by him were probably abnormal, their shape being perhaps due to accidental decapitation at the entrance of the boring and subsequent regrowth. The dimensions of the conidiophore and the conidia given by Giard are smaller than those obtained from the Kew specimens, but there is not much doubt that the latter belong to his species, which must now be known as *Hirsutella Barberi* (Giard).

125. *ISARIA TENUIPES* Peck

This species was described by Peck in the 31st Report, New York State Museum (1879), p. 44, as "Stem very slender, elongated, glabrous, lemon yellow, $1-1\frac{1}{2}$ inches high, divided above into a few irregular branches, which are wholly covered by the white mealy coating of conidia; conidia oblong-elliptical, $4-5\mu$ long. Dead pupae buried under fallen leaves. Center. Sept."

Pettit, in *Artificial Cultures of Entomogenous Fungi* (1895), described what he considered to be Peck's species, but as he stated that the conidia were oval to globose, $2.5-3.5\mu$, it would seem doubtful whether he had *Isaria tenuipes*.

I have not seen the type of *I. tenuipes*, but several examples of an *Isaria* which appears to answer to Peck's description were included in

a collection of entomogenous fungi submitted to me by the Mycological Herbarium, Cornell University. They were collected at Taughannock Gorge, New York, 3 September 1908 (No. 2906). The clavae are up to 2.5 cm. high, either simple, 0.3 mm. diameter below, with a narrow clavate head up to 1 mm. diameter, or branched above, the apices of the branches divided and flattened flabelliform, or, in the smaller forms, plumose. The colour is now white, the stalk becoming brownish in the larger specimens, but the ultimate branches have a yellow core when viewed by transmitted light. The stalk is pruinose, and the head is granular with minute spheres. These spheres are solitary at the ends of simple hyphae, $50-80\mu$ long, 2.5μ diameter, which arise singly from the clava, or as lateral branches of long, free hyphae. The apex of the hypha (conidiophore) bears a cluster of cells, either oval, $5 \times 4\mu$, or globose, about 3μ diameter, from which arise phialides, either broadly flask-shaped, $4-5 \times 1.5-3\mu$, or subglobose, 3μ diameter, with a conical neck. The conidia are hyaline, cylindric or oblong oval, sometimes inequilateral or subcymbiform, usually straight, sometimes slightly curved, $4-6 \times 1.5-2\mu$.

This *Isaria* belongs to the *Isaria japonica* group, a type of *Isaria* which is common in the tropics. In their fully developed form, the species of this group are much branched above, but the branches are somewhat distant and distinct from one another, so that the head is plumose, contrasting with the more solid head of the *I. farinosa* type. The long simple stalks of the conidiophores also add to the feathery appearance of the head. The conidia are usually cylindric, straight or curved. Lloyd's photograph of "*Isaria farinosa*" in *Synopsis of the Cordyceps of Australasia*, fig. 613, appears to be *I. tenuipes*.

This species should be compared with *I. furcata* Schw., *Synopsis N.A. Fungi*, No. 3067, if the type of the latter is available.

126. *ISARIA SUFFRUTICOSA* Cke. & Massee

Isaria suffruticosa was described by Cooke and Massee in *Grevillea*, xix (1890), 45, from a specimen on a hairy caterpillar collected at New England, Australia. Cooke gave a further account of it in *Vegetable Wasps and Plant Worms*, in which he stated that it was in tufts about 3 cm. high, and white. The stem was quite distinct, and either smooth or mealy, divided in the upper portion into very slender interwoven branches, which bore lateral branches up to the acute tips. The whole was "compacted of delicate threads, the extremities of which were free and bore the minute, ellipsoid conidia." The original description stated that the conidia are narrow-ellipsoid, $4-5 \times 1.5\mu$, borne singly at the apices of short sterigmata. A drawing which accompanies the type specimen shows a long hypha with short lateral branches bearing single conidia, but that was probably drawn from

a hypha to which conidia were merely adhering. The conidiophores on the specimen are quite different.

There is no insect in the type specimen now. It consists of a single clava, about 3 cm. high. The stalk is now brown, white pruinose, with numerous, slender, spreading branches in the uppermost centimetre. The core of the branches is yellow by transmitted light. The branches give off lateral branchlets, and both bear, as conidiophores, long, septate hyphae, 2μ diameter, which terminate in a cluster of globose cells, $3-5\mu$ diameter, bearing globose phialides with a conical neck, $4-6 \times 3-3.5\mu$, the whole forming a more or less globose head. The conidia are subcymbiform, or cylindric, or oblong-oval, straight or slightly curved, hyaline, continuous, $3-6 \times 1.5-2\mu$.

I. suffruticosa belongs to the *I. japonica* group. It may, indeed, be the same as *I. japonica*, but I have not seen an authentic specimen of the latter.

127. *ISARIA ONCOPTERAE* McAlp.

This species was described by McAlpine in *Proc. Roy. Soc. Victoria* (1894), p. 165. His account gives a few more details than are included in the description in Saccardo, *Sylloge Fungorum*, xi, 642.

The fungus occurred on the larva of *Oncoptera intricata* Walk. (Lepidoptera). "About twenty specimens were found near Melbourne, between August and October, inside the grassy tubes made by the larvae, and in every case either on a level with the surface or above it. All the infected larvae observed were nearly full-grown and dead, but in no instance were they found dead below the surface of the ground, although many tubes were examined, the larvae being always alive and apparently healthy when found below the surface." As many as fourteen clavae arose from one larva. The clavae were $1.8-3.6$ cm. high, at first purple, then brown ("dirty brown root colour") with a white apex, slender, branched, velvety, bearing conidia towards the apices of the branches. The conidia were spindle-shaped to oval, hyaline, $12 \times 6\mu$, borne on tips of hyphae at right angles to the clavae.

Apparently no specimens have reached England. From the description, the fungus appears to be a *Hymenostilbe*.

128. *SPICARIA (ISARIA) LAXA* Petch, n.sp.

This species was collected by Thaxter, on a wireworm at Kittery Point, Maine, September 1902 (Farlow Herb. 641). The larva, which was apparently buried, bears small, white, rudimentary clavae, or strands of mycelium, from all the sutures. These clavae are fasciculate, loose, linear, up to 2 mm. long, 0.1 mm. diameter, and consist of hyphae, 4μ diameter, which bear short, lateral, oval prophialides, 6μ long, 4.5μ diameter, with a group of phialides at the apex, or short,

stout, septate conidiophores, 4μ diameter, which bear whorls of three similar prophialides and a group of three divergent prophialides at the apex. The phialides are conical or flask-shaped, $6-11 \times 2-2.5\mu$, attenuated above into a slender sterigma. The conidia are fusoid, slightly curved, with one end acute, $5-6 \times 1\mu$.

The structure of this species is similar to that of *Isaria dubia* Delacroix. Indeed, it scarcely differs from the latter, except in the shape of the phialides. *I. dubia*, however, is the conidial stage of *Cordyceps gracilis* and occurs on Lepidopterous larvae, whereas *Isaria laxa* occurs on Coleopterous larvae and may possibly be related to *Cordyceps styphora*.

Spicaria (Isaria) laxa Petch, n.sp.—Clavis fasciculatis, laxis, linearibus, usque 2 mm. alt., 0.1 mm. diam., hyphis 4μ diam.; conidiophoris brevibus, crassis, 4μ diam., septatis, prophialides triverticillatos ferentibus; prophialidibus ovalibus, 6μ alt., 4.5μ diam.; phialidibus conicis vel ampullaceis, $6-11 \times 2-2.5\mu$; conidiis fusoideis, leniter curvatis, uno fine acutis, $5-6 \times 1\mu$. On Coleopterous larvae, North America.

129. *SPICARIA VELUTIFORMIS* Petch

This species, which occurs on spiders, was originally described from specimens from Ceylon. I have since examined specimens from the Hawaiian Islands, Palola Valley, Oahu, 3 April 1911 (Farlow Herb. 637) and from the Philippines, Stotsenberg, Pampanga, Luzon, 11-13 March 1923 (Farlow Herb. 6246).

130. *SYNNEMATUM JONESII* Speare

In *Mycologia*, XII (1920), 74, Speare described a new genus, *Synnematium*, with the species *S. Jonesii*. The fungus occurred on Hemiptera, *Mezira emarginata* and *M. lobata*, in Louisiana. *Synnematium*, in most respects, resembles *Tilachlidium*. It forms erect clavae, with lateral and terminal conidiophores which bear at their apices a globule of conidia embedded in mucus. The globules on the lateral conidiophores remain separate, but those on the conidiophores at the apex of the clava fuse into a large mass. The conidia are at first hyaline, but ultimately become brown or almost black. In addition, the fungus produces globose sclerotia, $150-200\mu$ diameter, white then brown.

Speare, no doubt because the conidia are cymbiform and embedded in mucus, compared his fungus to *Hirsutella*. This genus was originally described as a Basidiomycete, and in Saccardo, *Sylloge Fungorum*, it is still included in the Clavariaceae. Consequently in vol. XXIII of that work, *Synnematium* is also placed in the Clavariaceae. But both *Hirsutella* and *Synnematium* are Stilbaceae.

In the Farlow Herbarium, No. 6147, there is a specimen of *Synnematium Jonesii* on a beetle, *Philonthus* sp., collected by Thaxter at

Kittery Point, Maine, September 1893. The insect does not bear evidence of attachment to any substratum, but it is in a broken condition and that point cannot be determined with certainty. It bears numerous erect, stilboid synnemata, caespitose, simple, or divaricate above, with a white stalk and a black head. The stalk is about 0.5 mm. high, 40μ diameter below, expanding upwards to 80μ , and is composed of rather loose hyphae, the outer divergent and upwardly directed. The outer hyphae of the stalk are $3-4\mu$ diameter, septate, hyaline, verrucose, with free, clavate, verrucose tips. The basal hyphae on the insect are smooth, brown, septate, $3-5\mu$ diameter. From the stalk, long hyaline hyphae, 1μ diameter, diverge, and bear at the apex a minute globule of mucus, which contains broadly cymbiform conidia, $9 \times 4\mu$, with rounded ends, at first hyaline, becoming brown. Sometimes these lateral conidiophores are compound, i.e. the stalk consists of a small number of parallel hyphae. At the apex of the synnema, a cluster of conidiophores is produced, but all the globules of these conidiophores fuse into a depressed globose, apical mass, $90-120\mu$ diameter and $80-100\mu$ high, which turns black.

In lactic acid, the black heads are dark brown. They contain the same conidia as the lateral heads, irregularly arranged as in *Cephalosporium*, with some hyphae which are the entangled apices of the conidiophores. As in *Cephalosporium*, the conidia are produced apically and persist in a globule of mucus. The tenacity of the mucus in this fungus is remarkable, and it is difficult to separate the conidia. Both the conidia and the mucus turn brown. Though the conidia are, in general, fairly uniform in size, some of the globules on the lateral, simple conidiophores contain cymbiform conidia, $3-4 \times 1.5-2\mu$, or oval conidia, $3 \times 1.5\mu$.

Another specimen, Farlow Herb. 6262, on ? *Pardomis*, collected by Dr W. H. Weston Jr. at Washington, D.C. is in an earlier stage of development. The clavae bear lateral, simple conidiophores with small apical globules containing cymbiform conidia, but none of them has an apical mass of conidia. However, the larger clavae terminate in a loose apical pencil of conidiophores. In this specimen, the conidia are hyaline.

Both the specimens recorded above answer to the description of the conidial form of *Synnematium Jonesii*, but they do not bear any sclerotia. On the other hand the following specimens, which bore numerous sclerotia but no conidiophores or stilboid clavae, have been examined; on *Harpalus* (Carabidae), coll. G. F. Atkinson, Auburn, Cal., 8 August 1890 (Farlow Herb. 694); on a large leaf-hopper, coll. J. R. Johnston, Costa Rica (Farlow Herb. 6143); on *Proreus simulans*, Los Banos, Laguna, February 1932, G. O. Ocfemia; on a Membracid, *Basilides bipennis*, Njala, Sierra Leone, December 1931; on a beetle, *Promecotheca bicolor*, Fiji; on *Helopeltis*, Belgian Congo. In all

these specimens the sclerotia have the same structure. In lactic acid, when subjected to light pressure, they readily separate into their component cells, which are thick-walled, irregularly ovoid, $11-15 \times 8-10\mu$, or subglobose, 8μ diameter.

R. L. Steyaert and J. Vrydagh have published an account of the sclerotia on *Helopeltis* in *Étude sur une maladie grave du Cotonnier provoquée par les piqûres d'Helopeltis* (1933). They state that the sclerotium is composed of two kinds of cells, the inner hyaline and irregularly angular, the outer pale brown, more globose and smaller, both kinds having very thick walls. Grown in culture on "agar saccharosé et peptoné", the sclerotium produced a dense, white mycelium, on which sclerotia were again formed. These authors give the dimensions of the sclerotia on the insect as $48-100 \times 48-84\mu$ (mean 77.6×89.2), and in culture, $84-140 \times 84-128\mu$ (mean 100×117.4). The internal cells of the sclerotia in culture measured $14-24 \times 8-24\mu$.

Speare stated that in culture the sclerotia produced the synnemata. As Steyaert and Vrydagh failed to obtain the conidial form in culture, there seems to be some doubt whether the sclerotia on *Helopeltis* belong to *Synnematium Jonesii*. It would, however, appear advisable that all these sclerotia should be referred provisionally to that species.

131. *Aegerita insectorum* Petch, n.sp.

This species occurred on the larva of *Urophora solstitialis* L. (Diptera) in the flower heads of *Centaurea nigra* at the Entomological Field Station, Cambridge. It forms pulvinate sporodochia on the larva, but may also produce a white mycelium, bearing clusters of conidiophores, in the cavity occupied by the larva.

Aegerita insectorum Petch, n.sp.—Sporodochiis hemisphaericis, usque 0.3 mm. diam., 0.2 mm. alt., sordide albis; hyphis fasciculatis, ramosis, $4-7\mu$ diam., aut moniliformibus, segmentis ovoideis, aut subregularibus, segmentis oblongis medio contractis, $18-20\mu$ longis; conidiophoris irregulariter conoideis vel ampullaceis, $14-27 \times 5-9\mu$, interdum ovalibus, $12 \times 6\mu$, apice obtusis; conidiis hyalinis, levibus, globosis, $6-9\mu$ diam., vel ovalibus, $9-10 \times 5-8\mu$, apiculatis. On the larva of *Urophora solstitialis* L. (Diptera), Cambridge, England.

132. *Sporotrichum peteloti* (Vincens) Petch

In *Trans. Brit. Mycol. Soc.* xvi (1931), 55, it was pointed out that *Beauveria peteloti* Vincens was evidently a *Sporotrichum* parasitic on *Hirsutella Saussurei*, and in the same journal, xix (1935), 186, another *Sporotrichum* parasitic on entomogenous fungi, *S. columnare*, was described, differing from Vincens's fungus in the size and mode of attachment of its conidia. Since then several other specimens of *Sporotrichum* parasitic on entomogenous fungi have been examined, and it

is now possible to summarise the differences between the two species. Both produce white, conical or cylindric clavae, in addition to mycelium overrunning the insect, and both have more or less geniculate conidiophores. In *S. columnare*, the conidia are oblong-oval, one end acute, or pyriform, $6-11 \times 2-4\mu$, and are attached just above the angles of the conidiophore, sometimes in a group of two or three, as well as along the upper part of the conidiophore. In *S. Peteloti*, the conidia are smaller, oblong or oval, $3-5 \times 1-2\mu$, and are attached singly at the angles, or along the upper part of the conidiophore.

Revision of the specimens has shown that both species were represented in the collections enumerated in *Trans. Brit. Mycol. Soc.* xix (1935), 187, though the figure given there is that of *S. columnare*. The revised lists of hosts and localities is as follows.

Sporotrichum columnare. On a spider capsule, Maricao, 23 March 1916 (Cornell University, 730); on a spider capsule, Maraval Valley, Trinidad, coll. R. Thaxter, 18 February 1913 (Farlow Herb. 2578); on leaf-hoppers, Burbank, E. Tennessee, coll. R. Thaxter, 1896 (Farlow Herb. 6232).

Sporotrichum Peteloti. On *Hirsutella radiata* on flies, Essequibo, British Guiana, 11 November 1916 (C. B. Williams); on *H. entomophila* Pat. on beetles, Moruga, Trinidad, 14 January 1921 (W. Nowell); on *Cordyceps dipterigena*, Cranberry, N.C., July-August 1887, coll. R. Thaxter (Farlow Herb. 6221); on *C. dipterigena*, Burbank, Tennessee, 12 August 1896, coll. R. Thaxter (Farlow Herb. 6266); on a beetle, probably on *Hirsutella entomophila*, Trinidad, March 1917, coll. J. B. Rorer (Farlow Herb. 638); on razor-back hopper, Maraval Valley, Trinidad, 3 May 1913, coll. R. Thaxter (Farlow Herb. 2573).

133. *ACREMONIUM TENUIPES* Petch, nom.nov.

This species is represented in the Farlow Herbarium by two specimens, No. 728 and another unnumbered, on spiders, evidently parts of the same collection made by W. G. Farlow in a greenhouse at Cambridge, Mass., 24 January 1890.

The fungus forms a rather loose, large, white mass of mycelium, involving the body and the bases of the legs of the spider. The host was evidently lying free, unattached to any substratum. The hyphae are regular, up to 2μ diameter, and bear lateral conidiophores up to 20μ long, which are either subulate, 1μ diameter at the base, attenuated upwards, or very slender and hair-like throughout. The conidia are hyaline, oblong or oblong-oval, continuous, $2.5-5 \times 1.5\mu$, terminal and solitary on the conidiophores. The slender conidiophores appear to break up readily, as numerous short lengths are usually present in a preparation.

I have specimens of this fungus collected in England, on a spider's

egg cluster, Reffley Wood, near King's Lynn, 8 June 1930, and on a spider attached to a living leaf, Holt House Wood, near King's Lynn, 17 August 1931. Both these are in poor condition, and have been unrecorded, awaiting better material. In the former, the conidiophores are up to 55μ long, 1μ diameter below, attenuated upwards, and the conidia measure $2-3 \times 1.5\mu$. In the latter, the conidiophores are up to 66μ long, $2-2.5\mu$ diameter at the base, soon attenuated to a fine thread, and the conidia measure $2.5-3 \times 1.5-2\mu$.

Cavara, in *Fungi Longobardiae exsicc.* (1892), No. 240, issued specimens of a fungus on a spider, with a description and figure, under the name *Sporotrichum araneorum* Cav. Its mycelium was said to consist of very slender hyphae, $0.6-0.7\mu$ diameter, and its conidia were described as ellipsoidal, $2.5-3 \times 0.5\mu$, solitary at the apex of the branches. Cavara's figure shows a straight hypha which gives off lateral branches, some of which again bear lateral branches. These branches are about the same diameter as the main hypha at the base, but are soon attenuated into a long, slender thread. The conidia are solitary at the ends of the main hypha and the branches. The mycelium is shown covering the body of the spider in a large, cottony mass, and minute tufts emerge from the joints of the legs.

There is an example of Cavara, *Fung. Longob. exsicc.*, No. 240, in Herb. British Museum. The hyphae are $1.5-2\mu$ diameter, and bear the same conidiophores as described above for the American and British specimens. Cavara's measurement of the mycelium evidently refers only to the thinner part of the conidiophore. The conidia are oblong or oval, $2-5 \times 1-1.5\mu$. Perhaps a dropped figure in printing accounts for Cavara's smaller breadth. There is no doubt that the fungus is the same as the American and British species.

Sporotrichum araneorum is recorded by Ferraris, *Flor. Ital. Crypt.* 1, from Pavia, Sicily, and Verrua (Piedmont), while Lindau, *Rabh. Krypt.-Flora*, recorded it from Muskau in Silesia on the authority of Sydow.

Sartory, Sartory and Meyer, in *C.R. Soc. Biol.*, Paris, cvii, 14 (1931), pp. 53-55, gave an account of an unnamed species of *Verticillium*, which had been found in association with a *Sporotrichum* on spiders. Their description is from cultures. The conidiophores were said to be "droits", with several verticils of primary branches, $2-4\mu$ diameter. The uppermost primary branches were opposite. The lower primary branches bore verticils of secondary branches, $0.5-1.25\mu$ diameter. The branches bore apical, ovoid conidia, $3.5 \times 1.5\mu$, usually solitary, but sometimes in a group of three.

It seems most probable that this last-named fungus is the same as Cavara's. The slender dimensions of what are termed "secondary branches" and the size of the conidia point to that conclusion. But neither in Cavara's specimen, nor in the American and British ex-

amples, are the branches in verticils. They arise at irregular distances along either side of the hypha, as figured by Cavares.

As will be evident, the classification of this fungus has presented some difficulty. All observers are agreed that the conidia are usually solitary at the apex of the conidiophore or phialide, and consequently it cannot be placed in *Sporotrichum*. In this genus, the conidia are borne successively along the hypha, and though only the terminal conidium may be present, the scars of attachment of its predecessors persist.

As regards *Verticillium*, the alleged branches are not verticillate. Moreover, I have not observed erect, branched conidiophores in any specimen. Whether Sartory, Sartory and Meyer's statement that the conidiophores are "droits", means erect or straight is uncertain. The hyphae which bear the slender conidiophores or branches are certainly straight, but they are only part of the general mass of mycelium. The correct interpretation appears to be that the attenuated branches are simple conidiophores, lateral on repent mycelium. If so, the fungus is an *Acremonium*, and as the name *A. araneum* is already occupied, I propose to call it *A. tenuipes*.

Acremonium tenuipes Petch, nom.nov.; *Sporotrichum araneum* Cav., *Fung. Longob. exsicc.*, No. 240; *Verticillium* sp., Sartory, Sartory and Meyer in *C.R. Soc. Biol.*, Paris, cvii (1931), 53-55.—Mycelio albo, corpus et proximos articulos artuum pulvino albo tegente; hyphis regularibus, usque 2μ diam.; conidiophoris lateralibus, usque 66μ long., subulatis, basi $1-2.5\mu$ diam., mox attenuatis, vel ab initio capillaribus; conidiis hyalinis, continuis, oblongis vel oblongo-ovalibus, $2-5 \times 1-2\mu$, apicalibus, solitariis. On spiders, Italy, Germany, France, England, North America.

134. *ISARIA BRACHIATA* (Batsch) Schum.

This species is usually found on decaying fungi, though Saccardo, *Sylloge Fungorum*, iv, 589, added that it had been recorded on roots and leaves. Its colour is generally given as white. In September 1934 I gathered a specimen at Ingleton, Yorks., on a small mass of dead leaves; the main stem was pale brown. In October of the same year I found another specimen on a decaying agaric at West Runton, Norfolk, and in this the main stem was pale yellow when dry. The conidia in both specimens were oblong or narrow-oval, $3-6 \times 1\mu$. In Saccardo (*loc. cit.*) the conidia are given as ellipsoid, $3-4 \times 1.5-2\mu$. *Isaria brachiata* has *Cephalosporium* conidiophores and is a *Tilachlidium*. The type species of the latter genus, *T. pinnatum* Preuss, judging from the description, does not appear to differ from *Isaria brachiata*.

In the Farlow Herbarium, No. 759, there is a specimen of *I. brachiata* collected by Thaxter on an agaric, Cranberry, North Carolina, 1896. The conidia are oblong, $2-4 \times 0.75-1\mu$.

The object of the present note is to call attention to another mode of occurrence of *I. brachiata*. In the Farlow Herbarium there are several specimens in which *I. brachiata* is parasitic on entomogenous fungi. Sometimes the *Isaria* may arise directly from an insect which was probably first attacked by an entomogenous fungus, or it may appear on a *Cordyceps* or *Hirsutella* clava, sometimes covering it with a continuous conidial layer, so that the whole appears to be a conidial *Cordyceps* clava.

In No. 6234, on a leaf-hopper, collected by Thaxter at Burbank, East Tennessee, August 1896, the insect bears numerous clavae from all parts of the body. These are erect or repent, up to 1.5 cm. high, 0.15 mm. diameter, usually curved, linear, with a few short, lateral, perpendicular branches. They are covered with conidiophores, which are those of *Isaria brachiata*, with *Cephalosporium* heads containing narrow-oval or subcymbiform conidia, $2-4 \times 0.75-1\mu$. In No. 666, similar clavae, scattered or fasciculate, up to 8 mm. long, 0.25 mm. diameter, occur on a specimen of *Ophiocordyceps unilateralis*, collected by Thaxter at Kittery Point, Maine. No. 6227, "on Bombycid", collected by Thaxter at Burbank, 1887, contains a cocoon, apparently of *Apatela americana*, which bears two clavae, up to 3 cm. high, probably of *Hirsutella gigantea*, but these are covered by a uniform layer of conidiophores of *Isaria brachiata*. No. 6238, on Coleopterous larvae, collected by Thaxter at Burbank in 1896, contains two larvae, each bearing several immature *Cordyceps* clavae, up to 2.5 cm. high, which, in general, are linear, sometimes flattened above, and vary in colour from pale yellow to dark brown, but most of them have small white branches, closely adpressed to the clava, or small white tufts of mycelium, which give the clavae a mottled appearance; these small branches and tufts of mycelium are *Isaria brachiata*, though only one conidium was observed at the apex of the conidiophore in No. 6238.

I. intricata Fr. is said to differ from *I. brachiata* in having only a single conidium at the apex of the conidiophore. As the cluster of conidia in a *Cephalosporium* must begin with a single conidium, it is questionable whether that is a valid difference, if the structure of the conidiophore is the same. A specimen of *Isaria intricata* from Glamis, in Herb. Kew. ex Herb. Berkeley, has *Cephalosporium* heads, while a specimen of *Isaria brachiata*, Romell, *Fungi exsicc. praes. Scand.*, has some conidiophores with *Cephalosporium* heads and others with single apical conidia. On the evidence of these specimens, the two species are the same.

A REINVESTIGATION INTO THE CAUSE OF "BROWN OAK", *FISTULINA HEPATICA* (HUDS.) FR.

BY K. T. ST G. CARTWRIGHT, M.A., F.L.S.

(With Plates I-VII and 6 Text-figures)

THE heartwood of English oak (*Quercus pedunculata* Ehrh. and *Q. sessiliflora* Salisb.) is well known for its durability and for the beautiful figure which it gives when cut on the quarter.

The colour of the heartwood varies from a light brown to a deeper and warmer shade, but occasionally assumes a rich reddish brown (*Argus brown**) in some individual trees when it is known as "brown oak" (Pl. I, fig. 1 A). This abnormal coloration may be evenly distributed throughout the heartwood or may occur in bands running parallel to the grain, when it is often known as the tortoiseshell variety, or in the West Country as "lion oak".

"Brown oak" is much prized for use in decorative work, and therefore such timber in the past has commanded a high price.

As far as is known to me, no certain means of ascertaining whether the heartwood of a standing tree will show this coloration has been discovered, up to the present, except by taking trial borings with an auger.

A similar abnormal coloration of the heartwood occurs, though more rarely, in sweet chestnut (*Castanea sativa* Mill.) (Pl. I, fig. 2 A).

HISTORICAL

No investigation into the cause of "brown oak" had been published prior to Groom's paper (1), in which he concluded that:

(1) The abnormal coloration was due to the presence of a substance which gave the reactions of the ill-defined material termed "wound gum" or "wood gum".

(2) The production of this was due to a fungus which produced conidiophores of the *Penicillium* type and was able to utilize the tannin present in the heartwood.

(3) On certain specimens small sphaeroidal basidiocarps were produced which were identified by G. Massee as those of *Melanogaster variegatus* Tul. var. *Broomianus* Berk.; no connexion between the *Penicillium*-like fungus and the latter was, however, established.

* Colours printed in italics are from Ridgway's *Color Standards and Color Nomenclature*.

(4) The fungus was stated to cause a depletion in the tannin content.

The restriction of "brown oak" to the heartwood and the distribution of the dark coloration in the streaky type made it seem unlikely that a fungus of the *Penicillium* type could be responsible. On the other hand, while *Melanogaster variegatus* var. *Broomianus* might be suspected of having some mycorrhizal association, it was unlikely to be found in the heartwood of a tree. Therefore it was decided to make a further investigation into the cause of "brown oak".

MATERIAL

Specimens of "brown oak" were obtained during 1933 and the early part of 1934 from various sources.

(1) A 1 in. board in the green condition stated to have been cut from a tree grown in Sussex in the vicinity of Lewes.

(2) From Kimberley Park, near Norwich.

(3) Two small samples from the Dartington Hall sawmills:

(a) In this specimen the brown coloration was evenly distributed throughout.

(b) "Lion oak" showing distribution in longitudinal streaks.

(4) Hedgerow tree of *Quercus sessiliflora* Salisb. from Gooseacre Farm, Radley, near Abingdon, Berks.

EXAMINATION

Microtome sections were cut from the specimens, these being examined both in the unstained and stained condition. Unless otherwise stated, sections were stained by the picro aniline blue and safranin method⁽²⁾. Microscopic examination of unstained sections showed an abundance of golden to reddish brown material often in the form of globules, distributed mainly in the medullary rays and wood parenchyma (Pl. II, figs. 1, 2). Fungus hyphae could be observed with difficulty and did not appear to be abundant; their presence was usually associated with the discoloration (Pl. II, figs. 1, 2). In normal coloured wood no fungus could be detected as a rule. In stained sections it was possible to observe the distribution of the hyphae, although in the areas which were deeply coloured the hyphae remained unstained and could be detected only with difficulty. Thus there appears to be a definite association between the presence of mycelium and coloured heartwood. The hyphae are of very varied diameter, the large ones often being a deep honey colour, though some of the hyphae in the deeply coloured wood-parenchyma cells appear to lose their contents, thus showing in outline behind the dark contents of the cells (Pl. II, fig. 1). The finer active mycelium branches profusely, most frequently at right angles to the parent

hypha. These branches send out finer branches until in places a regular network of mycelium may be present. The fungus passes from cell to cell, mainly through the pits, but direct penetration through the cell wall also takes place, as is shown in Text-fig. 1. Secondary spores are rarely formed; they are usually terminal but may be intercalary.

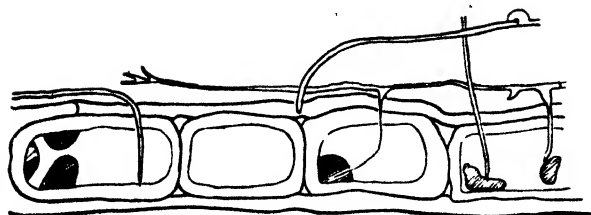
In addition to the profuse branching, the hyphae may describe loops. Clamp connexions, which are of the simple kind, are not abundant (Text-fig. 1). On staining sections of the wood with ferric chloride, the coloured areas assume a deep blackish blue tint, showing the presence of abundant tannin; but some of the reddish brown material does not assume this colour. Tests for the so-called "wound gum" gave no very definite results. The fact that the hyphae were frequently found investing the tannin, and that sometimes the tannin appeared to have become partially broken down and lighter in colour, suggests that the fungus obtains some of its nutritive material (presumably sugar) from this source. The so-called "wound gum", which does not give a tannin reaction, may be either an excretion product of the fungus or due to a direct degradation of the wood. More direct evidence for this has been obtained, and will be found on p. 77-82. Oil-like globules, which may be colourless or pale golden to reddish, are often present on the hyphal walls.

CULTURES

Cultures were obtained from all the specimens by means of a Pressler increment borer, plugs of wood being removed from the coloured areas. The outer portion of the plug was removed with a sterilized scalpel and the remainder partially buried in the medium. The medium on which the fungus was found to develop best was a 5 per cent Keppeler's malt agar which had been slightly acidified with malic acid; good growth was also obtained on 5 per cent malt agar. Development was also fairly good on a 2 per cent prune agar, but on 2 per cent malt agar growth was slow and weak.

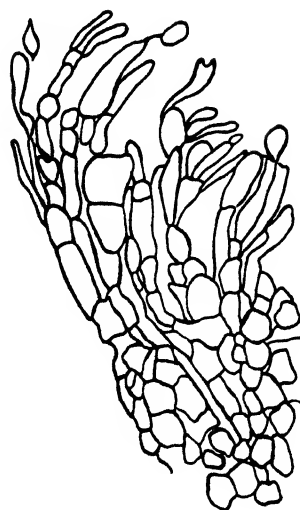
Growth was obtained from nearly all the plugs taken from the brown areas, though usually one to three weeks elapsed before it was evident, the length of time depending to a large extent on the period elapsing between felling of the tree and culturing; the fungus soon loses its vitality in the felled timber.

A few brown dusty cultures forming *Penicillium*-like conidiophores were obtained. This fungus was identified as one of the *P. divaricatum* group and is the same as that which has been obtained from oak timber showing yellow stain; it is probably synonymous with *Eidamia catenulata* Horne & Williamson, the fungus shown by Williamson (3) to be the cause of "golden oak". Cultures have been submitted to



0 10 20 30 40 μ

Fig. 1.



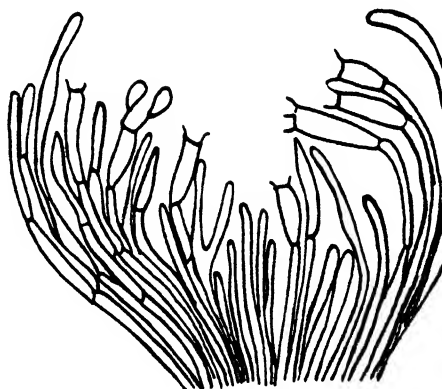
0 10 20 30 40 μ

Fig. 4.



0 10 20 30 40 μ

Fig. 2.



0 10 20 30 40 μ

Fig. 3.

Text-fig. 1. Portion of medullary ray cell in radial section illustrating penetration of hyphae through the walls. Note also tannin masses and a single simple clamp connexion.

Text-figs. 2 and 3. Section through portion of tongue-shaped sporophore developed on 5 per cent malt agar, isolated from "brown oak" in different stages of development.

Text-fig. 4. Ditto. One side of tube showing portion of hymenial surface in detail.

Mr E. W. Mason of the Imperial Mycological Institute, who confirms that it is *Penicillium divaricatum*, or one of this group; this, C. Thom⁽⁴⁾ states, is synonymous with *Poecilomyces Varioti*. It appears probable that this was the fungus which Groom isolated and considered to be the cause of "brown oak". The mycelium shows no resemblance to that observed in sections of "brown oak" and, moreover, the prevalence of this fungus in all kinds of timber being second only in frequency to that of *Trichoderma lignorum* and *Penicillium "glaucum"*, should ensure abundant "brown oak" if it were indeed the cause. Oakwood inoculated with this fungus becomes stained a golden yellow, but never assumes the deep brown tints typical of "brown oak".

Description of culture

At the beginning of growth, nearly colourless coarse mycelial threads issue from the wood plug. This mycelium forms a somewhat sparse tuft suggesting the early stage of such a mould as *Mucor* except for its very slow rate of development. In a few days this nearly colourless mycelium spreads on to the surface of the medium as a thin layer in which the individual threads can still be distinguished; as it becomes denser it becomes white. Growth is always slow, and the mat at this period resembles the texture of loose cotton-wool. Growth is more rapid in the dark; the optimum temperature lies between 25 and 27° C.

The centre of the mat is usually slightly depressed and slightly zoned, but the extent of this is variable. Some cultures remain white for several weeks, but usually the mat becomes *pale pinkish cinnamon* to *pale congo pink*, the centre being *straw yellow* which deepens to *mustard yellow*, the growing edge of the culture remaining colourless. Great variation in the colour development is shown in individual cultures even in a series of plates made from the same isolation, the range shading from almost white to *Mikado orange*, *orange-cinnamon*, *light vinaceous-fawn* and *vinaceous-fawn*, in addition to the pink shades given above. The coloration is shaded into zones, and tints of russet are often apparent. Pl. III shows a typical plate culture reproduced from a hand-coloured photograph and gives an accurate rendering of the main colour range. In spite of this wide variation the tints all lie in the same general range, and the whole appearance of the cultures renders identification easy and independent of microscopic examination.

In active cultures, drops of liquid are produced from the hyphae; these drops may be colourless when produced in cultures before colour has developed in the mycelium, but they are usually produced after coloration has developed; they are then pale amber to orange reddish or liver colour, resembling the juice abundant in an active

sporophore of *Fistulina hepatica*. These drops are acid, having a pH value of about 3.7. Secondary spores are usually formed in abundance on culture media.

In certain cultures, as they mature, sclerotium-like lumps are produced, these usually arising in the centre of the culture at the point of inoculation. Coloured drops similar to those on the mycelium are produced from these, and sometimes small tubes or plates indicating that these structures represent rudimentary sporophores (Pl. IV, fig. 1). Under the microscope, basidia, bearing spores averaging $5 \times 3 \mu$, have been observed; their formation is not confined to places where tube formation has commenced, instances having been found where the basidia are formed directly from the mycelium of the sclerotium-like lump.

In addition, tongue-shaped outgrowths are often produced, usually occurring in cultures grown in the dark. These may develop in as little as three days, in this respect their growth being more comparable with that of an agaric than of any *Polyporus* species so far cultured. Light appears to have a greater retarding action on the growth of the culture than is usual. The upper portion of these tongue-like structures is slightly swollen, so that they resemble a small species of *Phallus* in shape. On the swollen head rosette-like spots develop, this portion appearing like a zoophyte colony (Pl. IV, fig. 2). Sections show that the rosettes represent initial stages of tube formation, a normal hymenial surface bearing basidia and spores being formed. Two stages in tube development are shown in Text-figs. 2, 3, 4. The central portion upon which the tubes develop is composed mainly of large, loosely woven together conducting hyphae (Text-fig. 5).

Identification

The culture isolated from the sample plank of "brown oak" obtained from near Lewes, Sussex, did not lead to immediate identification, as at that time the wide variation in colour which occurs in different cultures was not fully appreciated. Microscopic examination of the mycelium revealed, however, the formation of secondary spores varying from 2 to 6μ or more in diameter. These were borne mostly on the aerial mycelium both terminal and intercalary, but were also observed more sparingly on the submerged hyphae (Text-fig. 6 and Pl. V, fig. 4).

The general appearance of the culture, and in particular its slow rate of growth, combined with the formation of secondary spores, suggested that the fungus isolated from the Lewes "brown oak" was *Fistulina hepatica*. Accordingly, comparative cultures were set up against a culture of *F. hepatica* already in the Forest Products Research Laboratory collection, obtained from sporophore tissue; the two were found to be identical. Further specimens of "brown oak" were

obtained from Kimberley Park, near Norwich, and in October 1933 two small specimens were obtained from the sawmills at Dartington Hall, near Totnes, South Devon. From these three specimens *F. hepatica* was also isolated from approximately 90 per cent of the plugs taken. Substantial evidence was thus obtained that *F. hepatica* is associated with the formation of "brown oak".

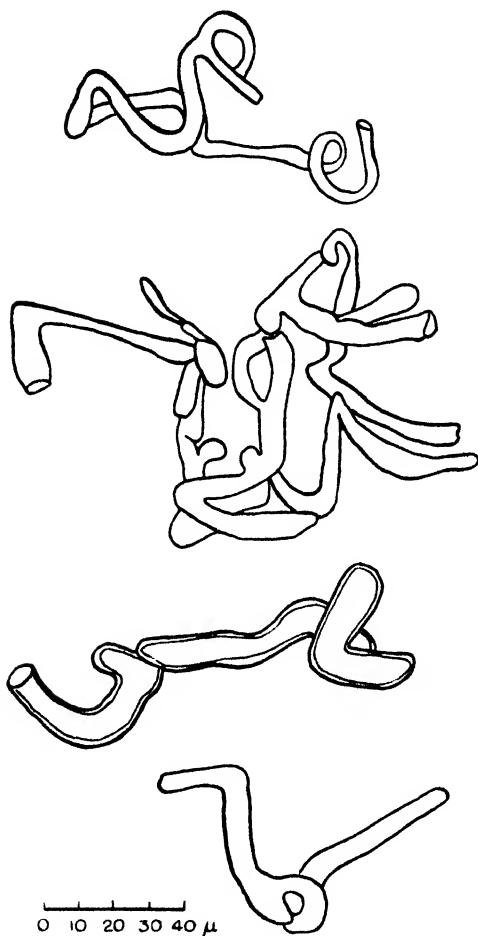


Fig. 5.

Text-fig. 5. Conducting hyphae from central portion of the same sporophore as figs. 2, 3 and 4.

Text-fig. 6. *Conidia* formed terminally on aerial hyphae.

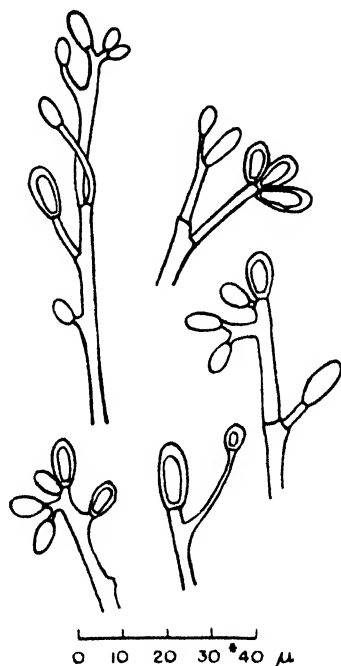


Fig. 6.

The fact that the brown coloration is said to occur usually towards the base of a tree or near the insertion of large branches, further supported the view that the fungus responsible for "brown oak" was of the heart-rot type, gaining entry through wounds occasioned by the fall of a branch old enough to expose heartwood, or through basal

wounds. The occurrence of "brown oak" is said to be more prevalent in mature trees, and this also falls into line with the above.

Conversion of sapwood into heartwood does not usually occur in oak until between the ages of twenty-five and forty years, according to conditions of growth, but at this period there would be only a small proportion of heartwood available, therefore the chance of exposure and consequent infection would be correspondingly less than in an older tree with a larger proportion of heartwood. The occurrence of "brown oak" in comparatively young trees, though it is rare, has occasionally been reported.

In addition to the evidence set forth above, *F. hepatica* is known to cause a red rot of oak timber, termed by the French "*pourriture rouge dure*" (5).

Blocks of oak sapwood and heartwood, sterilized by steaming on three consecutive days, were placed on an active culture of *F. hepatica* grown on malt-agar medium in special modified Kolle flasks. After six months they were removed from the flasks and samples of heartwood and sapwood from one flask examined. The wood had darkened and was distinctly of a richer colour than corresponding blocks which had been steamed but not exposed to fungus attack. Stained microtome sections showed a certain amount of mycelium (mostly confined to the large vessels) in the sapwood. In sections taken from the heartwood there was abundant fungus mycelium distributed throughout, and a certain amount of brown amorphous material was also observed. These experiments were somewhat inconclusive, and it is thought probable that, as in many of the heart rots affecting the standing tree, conditions required for the rather specialized metabolism of this fungus are not easily reproduced under laboratory conditions. Living trees have been infected with *F. hepatica*, and it will be possible to follow in them the development and rate of progress in the formation of "brown oak".

INOCULATION EXPERIMENTS

Inoculations were made by the removal of a plug of wood from an oak branch reaching into the heartwood and replacing it with one of "brown oak" containing *F. hepatica*. These inoculations were carried out in March 1934. The branch was amputated in November 1934, and the portion round the point of inoculation examined. Discoloration had taken place in both the sapwood and heartwood surrounding the inoculum, and mycelium of *F. hepatica* was observed in the wood at a distance of about 5 cm. from the inoculum (Pl. V). Some discoloration of the sapwood was probably caused by wounding and seepage of colouring matter, but the coloration in the heartwood resembled that of "brown oak".

It is proposed to follow the rate of progress of the fungus in relation to the spread of brown discoloration, but it will be some years before the results of the field experiments are completed.

Although the evidence appeared overwhelmingly to support the view that *F. hepatica* was the fungus responsible for "brown oak" formation, the failure to reproduce it in the laboratory made it necessary to apply a practical test. Accordingly, search was made for a suitable oak tree attacked by *F. hepatica*. This fungus is commonly observed on decrepit park trees which are often "stag-headed", and often almost completely hollow; but such a tree would obviously be unsuitable for selection, as complications due to the presence of other fungi able to cause decay would render any evidence obtained inconclusive.

A tree which appeared suitable for the purpose was found on the Radley College Estate, growing in a hedgerow. During the previous autumn a fruit body of *Fistulina* had been observed growing from the base of the tree. The tree, which was about ninety years old, although not a fine specimen, did not appear to be hollow, though some of the upper branches had been lost and it was slightly "stag-headed". Plugs were removed with a Pressler increment borer about two feet above the point at which the remains of the old fruit body could still be observed. The borings showed that the wood in this region was hard and was dark coloured. These plugs were placed in tubes of medium and sections were also cut from them. Sections showed the typical appearance of "brown oak" and *Fistulina* developed from the plugs placed in the culture tubes, thus showing that the fungus had spread at any rate sufficiently in the heartwood to give the desired information.

After this preliminary survey, the Radley College authorities were approached, and kindly gave me the tree. The tree was felled on 23 March 1934, being cut approximately one foot above the position of the fruit body. As felling proceeded and the heartwood became exposed, it was seen to be a rich reddish brown (Pl. V, fig. 3). The heartwood across the diameter was a uniform red-brown and this coloration extended up to the first branching and into the branches for a short distance where it began to run out into streaks as in "lion oak" (Pl. VI). The colour here was not so intense as in the main bole, tending to lessen in intensity after the first fork at eight feet from the butt end. The heartwood and sapwood appeared hard and sound throughout except where decay had entered through a dead branch, but this cone of decay did not proceed far and was not caused by *F. hepatica*. The "brown oak" was clearly defined in all the lower portion up to the first fork, extending right to the margin of the newly formed heartwood; beyond this a narrow band of heartwood of the normal colour could be detected until finally the colour ran

out in streaks. Beyond the area of brown in the portion where some of the heartwood was unaffected, a faint lilac colour could be detected in the wood adjoining the "brown oak", this being an area which the fungus was just invading. From the distribution of the "brown oak", it appears certain that infection originated through a wound at the base of the tree. As no figures are available for the rate of spread in a living tree, it is not possible to state with precision at what age the tree became infected. The slow growth of the fungus in the laboratory gives no reason for assuming that it grows at a similar rate in the standing tree. The extent and evenness of the coloration suggests that infection took place at an early stage when the tree was between thirty to fifty years old and that the fungus kept pace more or less with the conversion of sapwood into heartwood. The old scar marking the place of attachment of the sporophore suggests that this was the point of infection, and therefore it is possible that the spread of the fungus throughout the heartwood is comparatively rapid. Further data on this will be obtainable only when the results of inoculation experiments are available.

During the course of this investigation, samples of sweet chestnut were received for examination; these samples showed brown bands in the heartwood similar to those seen in the "brown oak" (Pl. IV, fig. 3). Sections showed the presence of fungus mycelium similar in appearance to that of *Fistulina* (Pl. IV, fig. 4). Cultures were made from this material and *Fistulina* obtained from it, thus showing that the fungus acts in a similar way in both oak and sweet chestnut.

PHYSIOLOGY

Tannin experiments

With a view to ascertaining whether *F. hepatica* is capable of utilizing tannin, an extract of oakwood which contained approximately 25 per cent tannin was obtained through the courtesy of the Forestal Land, Timber and Railways Co., Ltd. A series of concentrations of this extract was made up with 5 per cent Keppler's malt extract and 2 per cent agar; at the same time a similar series, omitting the malt extract, was also made up. In the series with malt, two plates containing 5 per cent malt and 2 per cent agar were included for comparison, as growth on this medium had been found to be excellent. In the series with tannin and agar but no malt, two plates with agar only were included. There were two plates at each concentration of tannin. The pH of the medium was difficult to arrive at by the colorimetric method, but lay between 3 and 4 in each, the addition of the tannin extract having little effect, though tending to

increase acidity. The results, averaged for each set of two plates, are summarized below:

Medium 2 % agar:

% tannin	0,	0.13,	0.25,	0.50,	0.75,	1.0,	1.25
Growth in mm. per diem	0,	2.8,	3.5,	3.6,	4.0,	4.1,	medium would not set

Medium 5 % malt and 2 % agar:

% tanin	0,	0.13,	0.25,	0.50,	0.75,	1.0,	1.25
Growth in mm. per diem	3.9,	3.9,	3.8,	4.2,	4.2,	4.2,	4.4

These results tend to show that *Fistulina* grows slightly better with the addition of oakwood-tannin extract and is able to obtain some of its nourishment from it, but as the tannin extract was not chemically pure, it does not prove that the fungus is necessarily breaking down the tannin. Further work on these lines is in progress. Pl. VII shows the stimulus to growth caused by the addition of tannin extract.

Attempts have been made to grow the fungus on a liquid tannin extract without agar, but although growth has been satisfactory for a time, it has not been considered sufficient to warrant chemical analysis of the medium before and after growth. From the fact that analyses of "brown oak" do not show much depletion in tannin and from the slowness of the action of the fungus on the mechanical strength, it is tentatively suggested that a degradation of the lignin and hemicellulose is taking place, leaving a brown gummy material as a degradation product.

As a matter of interest the following fungi were grown for comparison on the medium on which good growth was obtained with *F. hepatica*, namely 1 per cent tannin, 5 per cent malt and 2 per cent agar and using 5 per cent malt and 2 per cent agar as a control, repeating at the same time with *F. hepatica*. The results of the experiment are tabulated below:

	1 % tannin, 5 % malt and 2 % agar mm. per diem	No tannin mm. per diem
<i>Merulius lacrymans</i>	No growth	21.3
<i>Polystictus versicolor</i>	3.0	13.0
<i>Daedalea quercina</i>	1.5	3.8
<i>Polyporus sulphureus</i>	13.0	9.6
<i>Fistulina hepatica</i>	4.6	3.9

It will be seen that, apart from *Fistulina hepatica*, the only fungus of those tested showing stimulation of growth on the addition of the tannin extract is *Polyporus sulphureus*, which also occurs in the heartwood of oak. The effect of 1 per cent tannin on the other fungi is markedly to retard growth, as is illustrated in Pl. V, fig. 1, of *Polystictus versicolor* showing abnormal growth in a Petri dish. It is of interest to note that *Daedalea quercina* is a sapwood fungus. Growth was entirely

checked in *Merulius lacrymans* and this agrees with the fact that this fungus rarely grows on oak heartwood primarily, although under very bad conditions it can attack it.

Temperature and rate of growth

A series of Petri dishes using a 5 per cent malt agar with the addition of a small quantity of acid was inoculated from an active culture of *Fistulina hepatica*. The daily rate of increase in diameter at the different temperatures was measured, and the average daily rate of growth in millimetres at the different temperatures plotted. The figures were based on the average of ten plates. The results show that the fungus will grow over a wide range of temperature, growing appreciably at 9° C. and the optimum lying between 25 and 27° C. Growth dropped off rapidly above 30° C.

*The effect of *Fistulina hepatica* on the mechanical strength of oak*

It has been considered in the timber trade generally that "brown oak" is as strong as normal oak, although it is said that the "brown oak" is "milder". There was no reason to question this opinion prior to the present mycological investigation, and the results of Groom's work, in which he attributed the brown colour to the action of a mould type of fungus, strengthened this view.

The discovery that *F. hepatica* was the causal fungus suggested that mechanical tests should be carried out, as the Basidiomycetes are the group to which most of the active wood-destroying fungi belong, and in addition *Fistulina* is known to cause a heart rot of oak timber although, even in the late stages of decay, the wood never crumbles to a powder in the hand, as does wood attacked by a fungus such as *Polyporus sulphureus*. Accordingly the Timber Mechanics Section of the Forest Products Research Laboratory were asked to carry out mechanical tests.

Tests on a few small samples carried out at an early stage in the investigation had shown that there was apparently no appreciable loss in strength in the "brown oak", but as the history of these samples was imperfectly known and only a small amount of material was available, no conclusions could be based on them. More adequate material was available from the tree obtained from Gooseacre Farm, Radley, near Abingdon, although even here normal oak heartwood from the same samples could not be obtained in sufficient quantity for comparison, so that the results had to be compared with those from oak of the same specific gravity and correction made to allow for differences. The mechanical tests which were carried out were as follows

(1) Static bending test, which gives the normal strength and stiffness of timber under transverse loading applied without shock.

(2) Impact bending test, which records its toughness or resistance to shock.

(3) Hardness test, which measures the resistance of the timber to indentation.

(In the first preliminary experiment only static bending tests were carried out.)

These tests showed that "brown oak" was inferior in strength to normal oak of the same density. The "brown oak" was equal in density to timber from the lower range of oak which had been tested previously. The wood of the "brown oak" was markedly softer and more brittle in comparison with normal oak of equal specific gravity. The decrease in strength ranged from 11 per cent in static bending at 12 per cent moisture content to 32 per cent in hardness when in the green condition. These results explain the fact that the wood of "brown oak" is milder and therefore easier to work than the wood of normal oak. There is no need to apprehend the continuation of decay after felling and drying, and "brown oak" can safely be used for decorative purposes where great strength is not required.

A full account of this side of the investigation, with figures obtained from the tests, has already been published (6), and it is hoped that later it may be possible to carry out further tests on material obtained from field inoculation experiments, in order to gain an insight into the rate of progression of attack in relation to alteration in mechanical strength and to correlate this with chemical alteration in the wood, as has been done in Sitka spruce attacked by *Trametes serialis* Fr. and in ash attacked by *Polyporus hispidus* (Bull) Fr. The second series of tests shows that although the strength is not greatly impaired in certain respects, there is a very definite softening of the wood and the timber becomes "brash". The absence of any appreciable loss in strength in the first series can be accounted for mainly because the action of the fungus is probably not rapid.

Estimations of tannin content show a decrease of tannin comparable with that found by Groom; but the brown material does not appear to be a tannin, as on extraction little or no alteration in the colour has been observed.

It has not yet been found possible to extract the brown material for analysis, as it is insoluble in all the commonly employed reagents. Further work is in progress with the object of elucidating the chemical action of the fungus, which appears to be somewhat unusual. The fungus, as is shown by mechanical tests, acts on the cell walls of the wood, but as its action is not rapid and does not cause much loss in strength in the early stages, it appears unlikely that the cellulose is acted upon to any great extent initially, and it is suggested that the action is probably on the tannin, on the lignin constituents of the wall and on the pectic substances (possibly the hemicelluloses), the

brown substance representing the residue. Such action probably would not cause much loss in mechanical strength, though it might soften the timber.

GENERAL DISCUSSION

In the tree from near Radley there was very little sign of decay although, judging from the extent of "brown oak" formation, the fungus must have been present for many years. Examination of the fungus in the wood shows that, apart from occasional bore holes, there is little sign of cell-wall destruction such as cross cracking or general thinning, as is usually found in wood attacked by an active wood-destroying fungus. These facts, in combination with the presence of the deep colouring matter in the cells, point to a somewhat unusual type of metabolism.

The two most common hosts of *Fistulina hepatica* are oak and sweet chestnut, both of which contain abundant tannin; the number of species of fungus which attack these two trees is limited, further suggesting a somewhat specialized metabolism. This may be an acquired toleration to tannin as is possibly so with forms which produce serious decay in oak, such as *Polyporus sulphureus*; or it may be, not so much that tannin acts as a toxic agent as that it reduces the amount of oxygen available and that such fungi require less oxygen. The fact that the dense timber of yew (*Taxus baccata*), is readily attacked by *Polyporus sulphureus*, supports this view. A third possibility is that some fungi may be able to obtain part of their food material by breaking down the tannin and obtaining sugars from it. This appears to be the most likely hypothesis for *Fistulina hepatica*, which is apparently able to obtain its food material from the heartwood for a long time without affecting the cell wall of its host. The biology of this fungus should afford an interesting study for the forest mycologist in co-operation with the biochemist.

The work of Dr R. C. Fisher, entomologist at the Forest Products Research Laboratory, shows that wood partially decayed by *Phellinus cryptarum* Karst. is preferred by the Death Watch beetle to sound wood. It is possible that "brown oak" may prove to be more liable to Death Watch attack, but so far no definite evidence on this point has been obtained.

SUMMARY

1. *Fistulina hepatica* has been shown to be the cause of "brown oak", and of a similar discoloration in sweet chestnut (*Castanea sativa*).
2. The cultural characteristics of the fungus are described.
3. A description of the fungus in wood and of its behaviour in culture are given.

4. The optimum temperature for growth in culture was found to lie between 25 and 27° C.

5. Preliminary experiments on the metabolism of *Fistulina hepatica* indicate that it is in some respects different from that of a normal wood-destroying fungus causing a brown rot: the fungus may obtain a portion of its sugar supply from the tannin. It is suggested that the brown amorphous material may be a degradation product of the lignin and hemicellulose brought about by the action of the fungus.

6. Results of mechanical tests (6) show that no appreciable loss in strength is caused in the early stages by the action of *F. hepatica*, but later considerable softening takes place and the timber becomes somewhat brittle.

7. There is no risk of trouble being caused by the continuation of decay in the converted timber, and the fact that "brown oak" is easier to work than normal oak is an advantage where great strength is not required.

To W. A. Robertson, Esq., Director, Forest Products Research Laboratory, thanks are due for permission to publish this paper.

In connexion with the provision of material, assistance was kindly given by Radley College Estate and Dartington Hall, Ltd.

Grateful acknowledgement is also made to my colleague, W. G. Campbell, Esq., for assistance with the chemical work, and to W. R. Hutchins, Esq., for the photography.

EXPLANATION OF PLATES I-VII

PLATE I

Fig. 1. Illustrations of brown (A) and normal (B) oak.

Fig. 2. Illustrations of brown (A) and normal (B) chestnut.

PLATE II

Fig. 1. Photomicrograph of tangential longitudinal section showing reddish brown material in medullary rays and wood parenchyma. $\times 250$.

Fig. 2. Ditto. Radial longitudinal section. $\times 375$.

PLATE III

Four-week-old culture of *Fistulina hepatica* isolated from "brown oak" growing on 5 per cent malt agar acidified with malic acid.

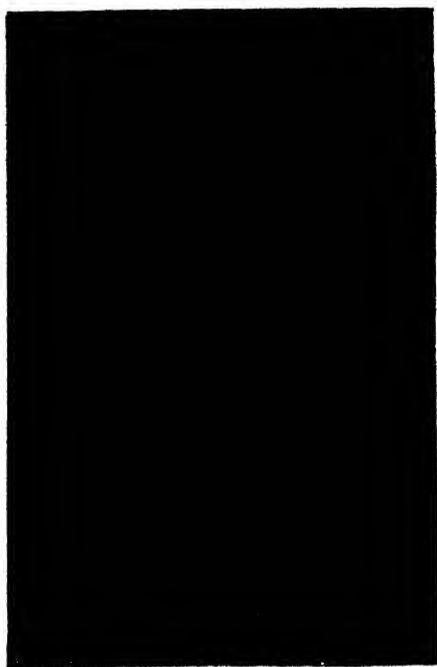
PLATE IV

Fig. 1. Incipient sporophore formed at point of inoculation on 5 per cent malt-agar medium after 2½ months.

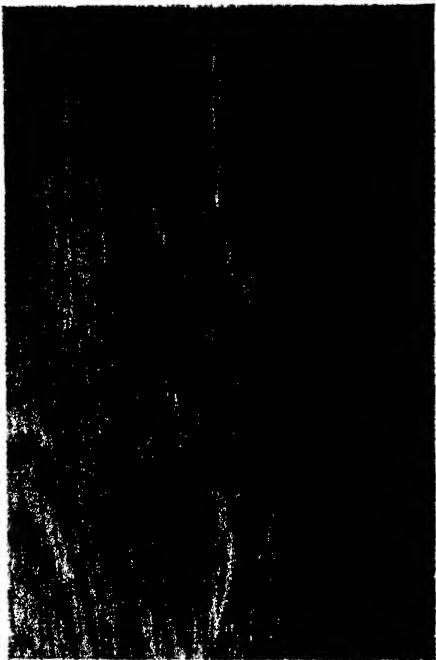
Fig. 2. Tongue-shaped sporophore developed on 5 per cent malt agar in the dark.

Fig. 3. Sweet chestnut showing brown coloration caused by *F. hepatica*.

Fig. 4. Photomicrograph of radial longitudinal section of wood of sweet chestnut cut across brown-coloured heart and showing hyphae of *F. hepatica* in the medullary-ray cells $\times 225$.

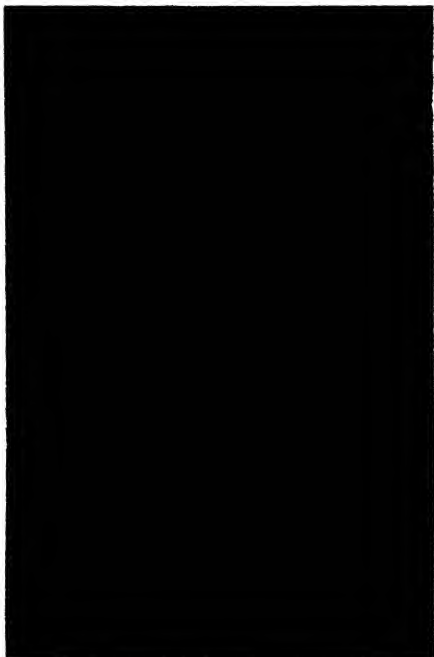


A

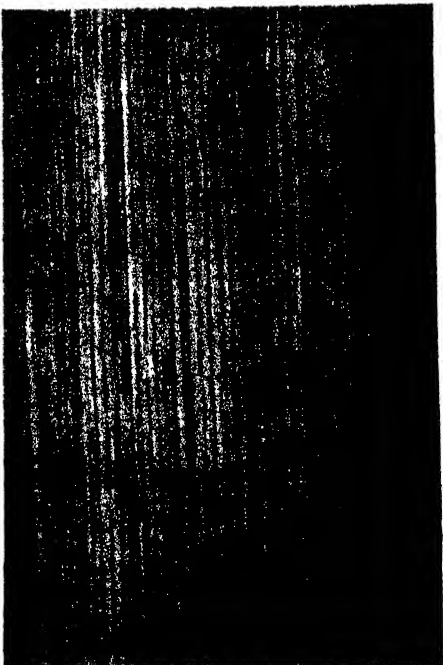


B

Fig. 1



A



B

Fig. 2

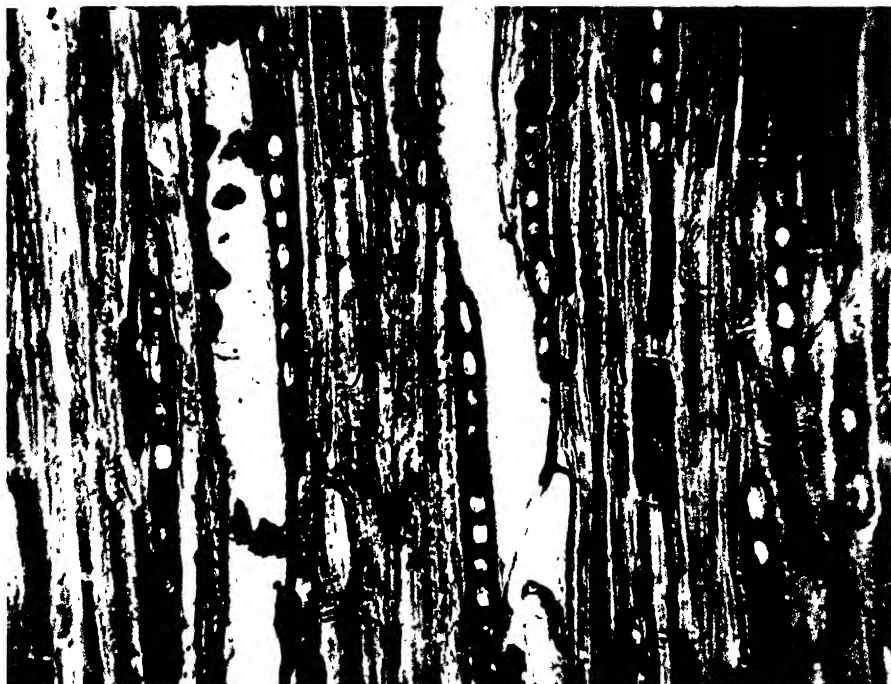
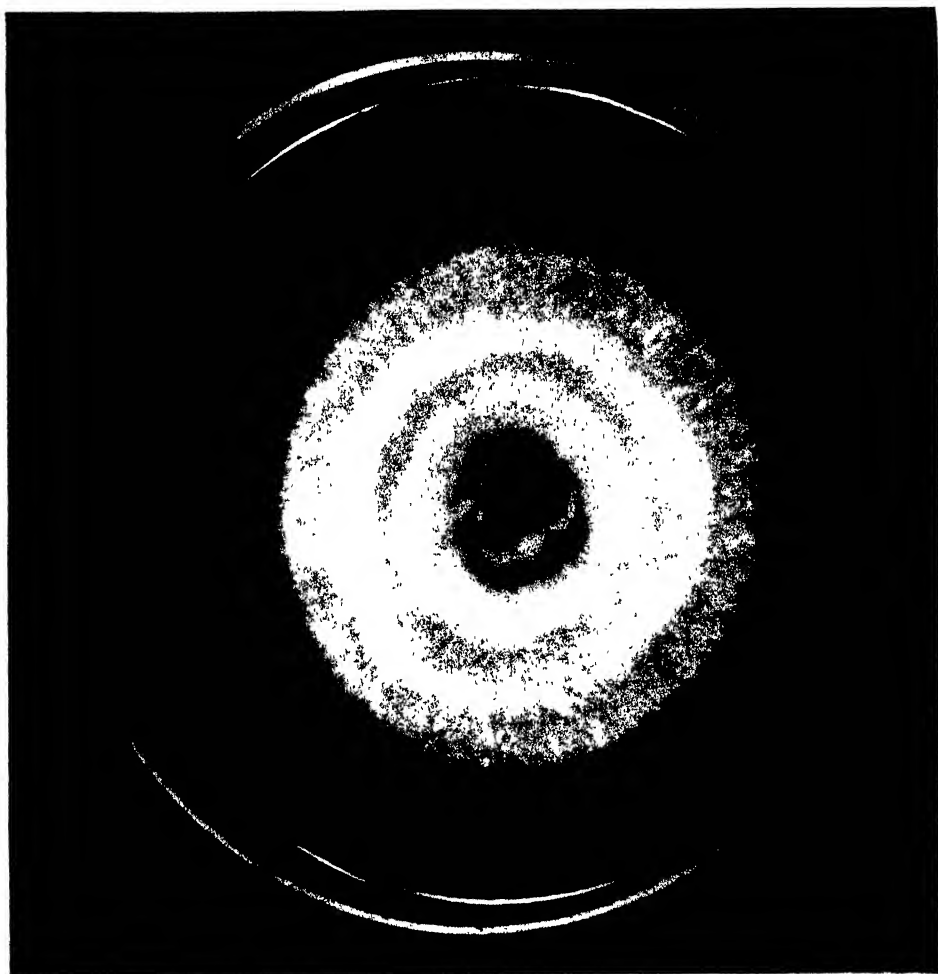


fig. 1



Fig. 2



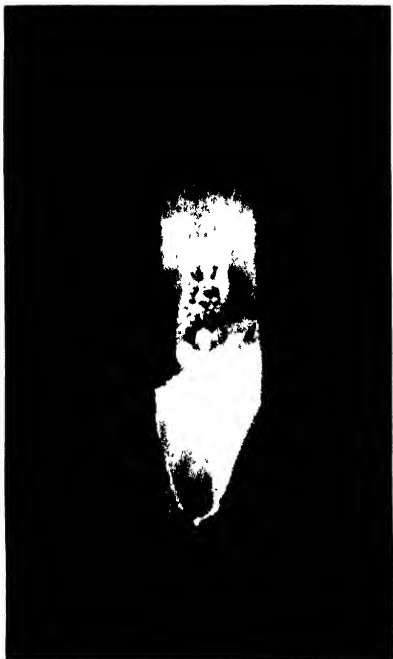


Fig. 1

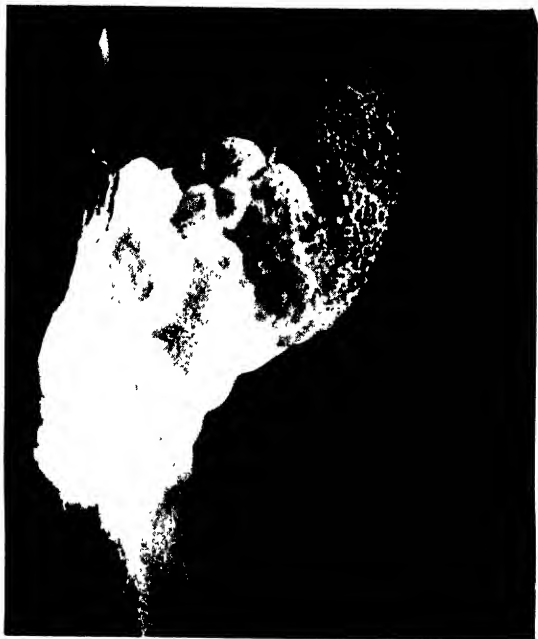


Fig. 2

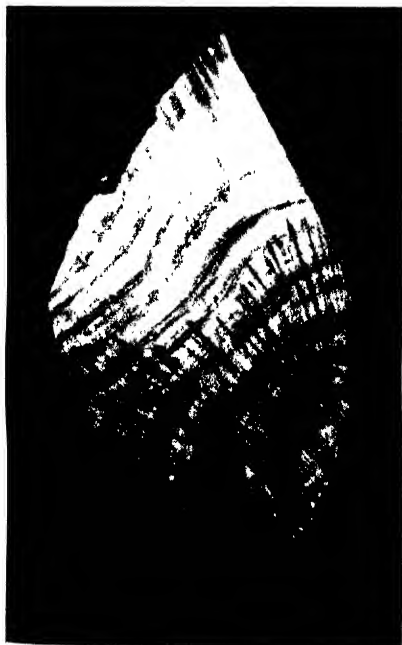


Fig. 3

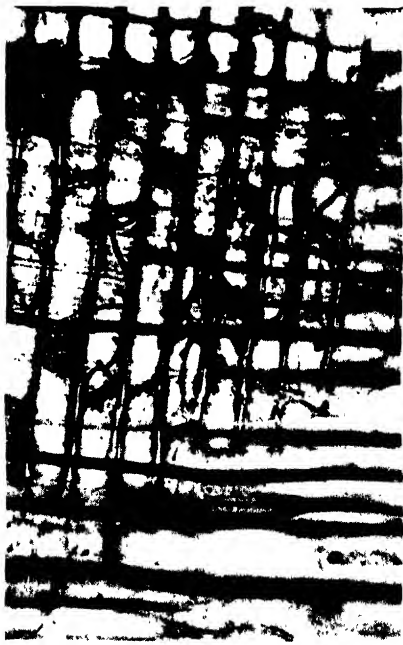


Fig. 4

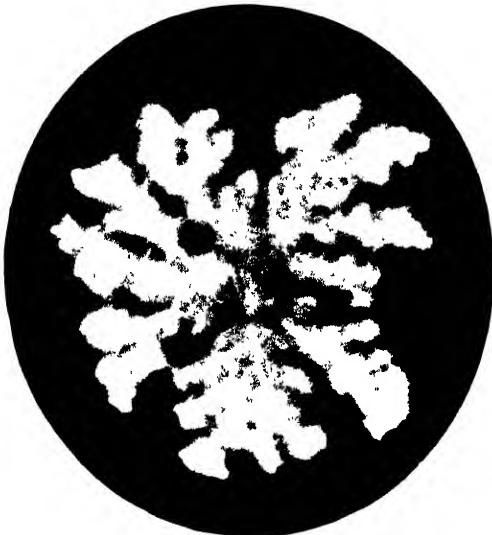


Fig. 1



Fig. 2



Fig. 3



Fig. 4





1.25 per cent tannin
2.00 per cent agar
(medium sloppy)



1.25 per cent tannin
5.00 per cent malt
2.00 per cent agar



1.0 per cent tannin
2.0 per cent agar



1.0 per cent tannin
5.0 per cent malt
2.0 per cent agar



Control
2 per cent agar



Control
5 per cent malt 2 per cent agar

PLATE V

- Fig. 1. *Polystictus versicolor* showing abnormal growth after ten weeks on malt-agar medium to which 1 per cent tannin extract had been added.
 Fig. 2. Portion of oak branch inoculated by means of Pressler increment borer. After seven months, showing discoloration caused by *F. hepatica*.
 Fig. 3. Oak tree from Radley at commencement of felling operations showing brown heart.
 Fig. 4. Photomicrograph showing secondary spores formed in culture. $\times 400$.

PLATE VI

Top portion of plank from Radley oak tree showing distribution of the brown heart at the upper extent to which the infection has reached. Note that in this area the brown coloration follows the outer limit of the heartwood on the left but that on the right normal coloured heartwood is also present.

PLATE VII

Petri-dish cultures showing stimulation of growth by the addition of oak wood tannin extract.

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FURTHER OBSERVATIONS ON THE SYSTEMIC INFECTION OF *LOLIUM*

By KATHLEEN SAMPSON, M.Sc.

(With Plates VIII and IX and 20 Text-figures)

I. A SECOND ENDOPHYTIC FUNGUS IN *LOLIUM PERENNE*

IN a previous study of the endophytic fungus which frequently occurs in certain species of *Lolium*, pedigree cultures of *L. perenne* were kept under observation for a number of years. The microscopic examination of pith scraped from a flowering stem and stained with cotton blue offered a simple and efficient method of detecting infected individuals in a mixed population⁽¹⁶⁾. One of the parents selected in 1929 at the inception of the experiment was noted as containing a mycelium which retained only a little of the blue stain. As the work progressed, this "faint" type of mycelium was recognized repeatedly and without difficulty and noted as such in the records. Subsequent results extending over a period of six years have shown that it may be transmitted either by seed or by vegetative propagation of the host in which it produces no obvious symptoms of disease. To this extent it resembles what may be termed conveniently the first *Lolium* fungus. It differs from this organism, however, in certain mycelial characteristics, in its failure to form a thick hyphal layer in the seed, and above all in the fact that it can readily be brought into culture on agar media. Unfortunately, apart from the production of microconidia, it has remained persistently sterile, and* like the original *Lolium* endophyte it has still no certain taxonomic position. The differences between the two types of infection in *Lolium* and in the invading fungi themselves are, however, of such a nature that it seems at present more reasonable to regard them as distinct organisms than to suggest any direct relationship between them. Evidence for this view will appear in the description which follows. Both types are also considered to be distinct from *Epichloe typhina*, which causes a similar systemic and latent invasion of *Festuca rubra*⁽¹⁵⁾.* It appears, therefore, that at least three fungi may lead a barren and seemingly

* Since the work on *Epichloe typhina* was published, further evidence of latent infection came accidentally to my notice. A plant of *Festuca rubra* which had been used successfully for a number of years by Dr Jenkin in his genetical work was found to carry mycelium which proved to be *Epichloe typhina*. The fungus caused no appreciable damage and never fruited in clonal lines of this plant.

innocuous existence in association with members of Gramineae. *Festuca* and *Lolium* are somewhat closely related but we have no evidence as yet that there is any near affinity between the invading fungi.

Transmission by seed and propagants

The experimental work has been carried out with pedigree cultures of plants originating in a single individual (No. 1972) selected in 1929. Since open pollination was permitted, the maternal parent alone could be recorded. Parallel with the progeny trials were various series of propagants obtained by the vegetative division in autumn of plants from which seed had been taken in the previous summer. The data collected over a period of six years are summarized in Table I. In column A appear the results from plants derived

Table I. *Pedigree populations of Lolium perenne* (No. 1972) selected 1929 as containing "faint" mycelium

Year		A. Parent infected. Mycelium in flowering stem		B. Parent fungus free. Mycelium in flowering stem	
		Present	Absent	Present	Absent
1930	Progeny	4	6	—	—
1931	Progeny	6	14	—	—
1932	Progeny	9	6	0	10
1933	Progeny	60	22	0	60
	Propagants	20	0	0	12
1934	Progeny	17	15	0	10
	Propagants	22	7	0	12
1935	Propagants	8	3	0	7

directly from parents which were known to be carriers of the "faint" type of mycelium. In column B are recorded the observations made upon the progeny and propagants of certain fungus-free plants which appeared from time to time in the cultures. The relatively large number of negative results recorded in the infected series (column A) is due to two reasons. In the first place, the faint mycelium was not so easy to trace in a pith-scraping as the deeply stained mycelium of the first endophyte (cf. (16), Table III, p. 341) and was undoubtedly overlooked sometimes. Such errors were detected by the examination of additional stems from particular plants and by observations on propagants in a subsequent season. The same procedure proved that negative results were sometimes due to the failure of the fungus to penetrate every part of a plant and the consequent appearance of fungus-free individuals in the pedigree lines. When such plants were multiplied by seed and by division they remained non-infected during an experimental period of four years (Table I, column B). Incidentally it may be noted that parallel cultures of *L. perenne* (No. 1975) derived from a plant selected as fungus-free in 1929 also remained true for

this character during the complete period and the same holds good for the lines of *L. perenne* (No. 1971) which escaped infection by the first endophyte ((16), p. 341). The origin of fungus-free races was rarer in the latter type of infection than it was in the second endophyte now under discussion. There is no reason to think that either type of infection came from the soil.

Although transmission of the present fungus by seed does not proceed without exception, a high proportion (over 70 per cent) of infected individuals does occur in the families which come directly from an infected female plant. Vegetative propagation is evidently even more efficient as a means of disseminating the fungus, since transmission occurred with one possible exception in all the propagants taken from infected plants.

During the period of this experiment, which covers five generations of the host, the characteristics of the mycelium and the type of infection have shown no sign of change. Fructification of the fungus on the host has been sought for in vain.

In view of the comparative ease with which fungus-free races can be obtained in cultures of rye-grass containing this particular endophyte, it might seem a simple matter to estimate the effect of the fungus upon the general growth of the plant. That this cannot easily be done arises from the fact that *L. perenne* is a cross-pollinated species and therefore markedly variable in both vegetative and floral characters. All that can be said is that the presence of the fungus is not associated with necrosis of tissue or with any visible pathological symptoms. Possibly more exact information concerning its effect on the host may be forthcoming from a study of the growth of reciprocal crosses between infected and non-infected plants.

Characteristics of the mycelium and its distribution in the plant

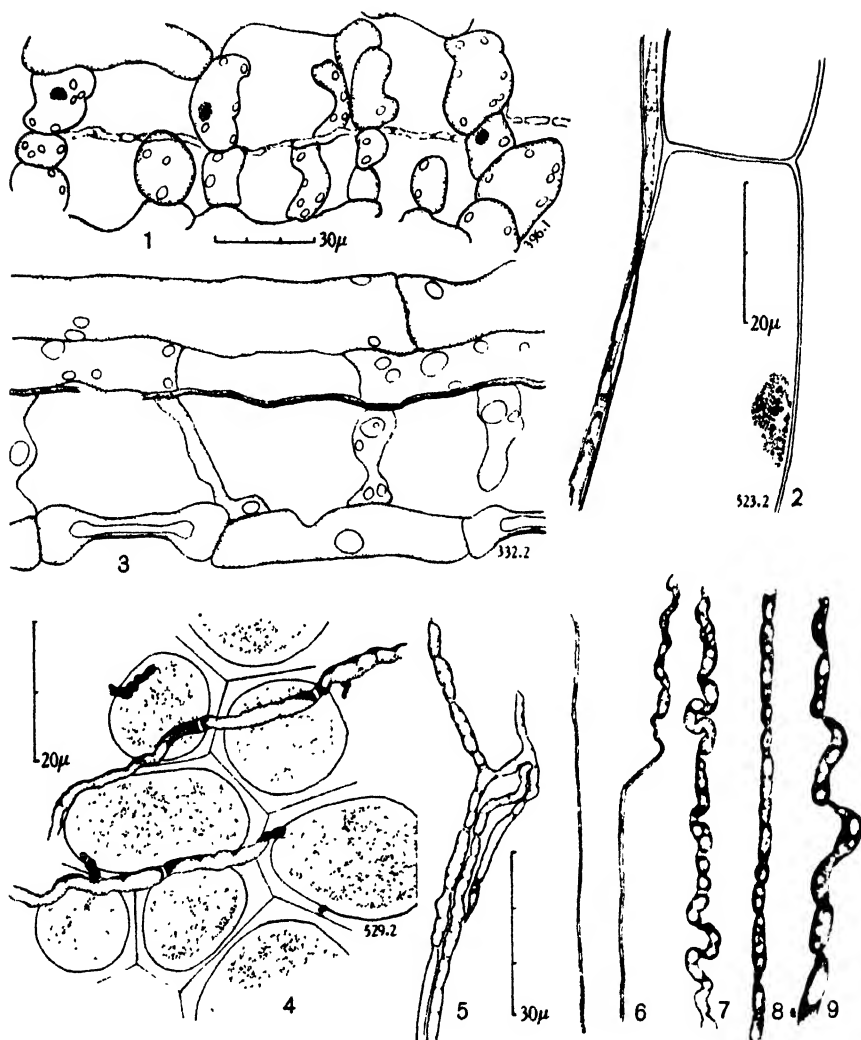
Mycelium is usually present in abundance in the pith scraped from the stem of an infected plant carrying the second endophyte of *Lolium* but it absorbs comparatively little of a protoplasmic stain, making a "faint" preparation in contrast to the richly coloured mycelium in a stained mount of the first endophyte. The differences seem to rest chiefly on the fact that the cells of the former have large vacuoles and sparse cytoplasm, while the latter is only slightly vacuolate and the contents of the cells look dense. This contrast is so marked that the two types can be distinguished with a high degree of accuracy (Pl. VIII, figs. 1, 3, and Text-figs. 5, 8, 9). Both fungi have rather closely septate mycelium but in the second endophyte the branching is more frequent and parallel strands or even wefts of mycelium are more often seen in pith preparations. The mycelium of the original endophyte, whether it is in *L. perenne*, *L. temulentum* or

L. remotum, usually occurs in the pith as long separate strands, often so much constricted at the numerous septa that a chain-like effect is produced. This is enhanced by a slight contortion which appears to result from elongation occurring in a limited space. A similar effect is sometimes produced in the mycelium of *Epichloe typhina*, but normally this fungus is characterized by the exceedingly fine straight lengths of mycelium which can usually be distinguished in pith mounts from both the fungi of *Lolium* (Pl. VIII, fig. 2 and Text-figs. 6, 7). In microtomed sections of leaves, mycelia of *Epichloe typhina* and the first *Lolium* endophyte look almost exactly similar.

Experimental evidence for the transmission of the second endophyte by seed and by vegetative propagation led to a search for the fungus in different parts of the plant. Portions of root, tiller buds, stems, ovules and ripe caryopses were fixed, embedded and stained with gentian violet by the methods previously used with *Epichloe typhina* (15). For comparison, material of *Lolium perenne* containing the true endophyte was likewise fixed and studied.

Mycelium of both the fungi of *Lolium* is intercellular. The second type, which absorbs little stain, is not nearly so easy to trace in the different tissues as the first, but both have been found widely distributed in the tiller buds of infected plants in both summer and winter. Text-figs 1, 3 show sections of young leaves from two infected plants with the characteristic mycelium of their particular endophytes crossing the intercellular spaces of the spongy parenchyma. I have not found either fungus in the root, but the difficulties in detecting it there are considerable and consequently little reliance can be placed on a negative result.

Passage up the flowering stem has been mentioned above. The pith in particular carries many parallel strands of mycelium. Invasion of the ovule and seed by the true endophyte has been described by other authors (5, 7). The most characteristic feature of this fungus seems to be the thick zone of mycelium which it forms in the ripe caryopsis outside the aleurone layer (Pl. IX, fig. 7). Such a zone has never been found in fruits containing the second endophyte. There is no doubt, however, that invasion of the flower occurs. Mycelium of the "faint" type has been seen in sections of the lodicules, at the base of the young ovary and in several mature fruits. In Pl. VIII, fig. 6 and Text-fig. 4 it is shown lying over some aleurone cells which are seen in surface view. In the ripe fruit it has been found only in this position, never in the embryo, but this may be due to the difficulty in detecting it among closely packed cells. This difficulty makes it impossible to decide by microscopic methods how many fruits do or do not carry this fungus. Data from the pedigree cultures indicate that while examples of failure are certainly not rare, seed transmission occurs in a high percentage of cases.



Text-figs. 1-9

1. Leaf of *Lolium perenne* (No. 1972) collected in February 1933 and cut in longitudinal section, showing the faintly stained mycelium of the second endophyte traversing the spongy parenchyma.
2. Mycelium of the same fungus in the leaf sheath of a young shoot collected in July. The mycelium is highly vacuolate like that taken from pith (Fig. 5).
3. Leaf of *Lolium perenne* (No. 1971) in longitudinal section showing the deeply stained mycelium of the first endophyte. It is scarcely distinguishable from that of *Epichloe typhina* in leaves of various grasses (15, Pl. VIII, fig. 1).
4. Mycelium of the second endophyte lying over some aleurone cells in a ripe caryopsis of *Lolium perenne*.
- 5-9. Mycelia from pith scrapings stained with cotton blue. 5. Branched vacuolate mycelium of the second endophyte of *Lolium perenne*. 6, 7. *Epichloe typhina* from a healthy inflorescence of *Dactylis glomerata*, showing the typical slender form grading into a coarser type which more nearly resembles that of the *Lolium* endophyte (8, 9).

Isolation and growth in culture of the second endophyte

It is not difficult to remove long strips of pith from grass stems and to place them on solidified agar without serious contamination. Flowering stems from plants carrying *Epichloe typhina* treated in this way yielded pure cultures of that fungus⁽¹⁵⁾. In May 1933 experiments were set up to see if this technique might provide a method of bringing into culture either of the *Lolium* fungi. Immediate success was obtained with the second endophyte. Strips of pith laid in close contact with clear nutrient agar were examined microscopically at intervals and the mycelial strands present were seen to branch, putting out rather fine, straight hyphae which finally formed a silky-looking colony. It was not unusual to obtain a pure culture immediately, all the pieces of pith on a plate giving the same type of mycelial growth. Contaminations which occurred on some plates were easy to detect.

In 1933, twenty-seven separate isolations, taken from three distinct plants, were kept in culture for a time but they showed no noteworthy differences. Fresh isolations made in 1934 and in 1935 agreed with stock cultures from the first isolation experiments. Since all the material originated from one plant, it is probable that all isolants represent one strain of the fungus.

Results of inoculation experiments are not available,* but there is little doubt that the cultures represent the "faint" type of mycelium in the pith, since the initial stages of growth of this mycelium have been traced under the microscope and the same fungus has been isolated many times over a period of three years. Moreover, plants lacking the fungus were tested occasionally by placing strips of their pith on agar but this remained free from mycelial growth.

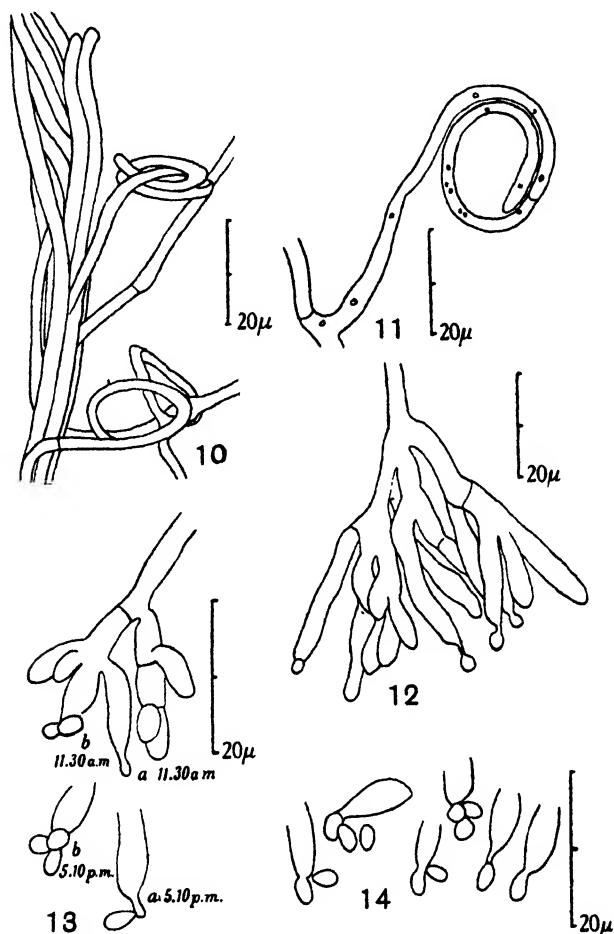
The fungus grows readily but somewhat slowly on a wide range of substrata, producing on agar media rather compact fluffy aerial mycelium, white to cream colour, in appearance like a miniature powder puff, while the submerged mycelium is often accompanied by a chrome-yellow stain, especially striking on oatmeal agar. Under a lens the aerial mycelium is seen to be funiculose, especially when lying near glass.

In branching and vacuolation the mycelium from cultures is not unlike that taken direct from the pith of the plant. In addition it produces rather numerous spirals (Text-figs. 10, 11), especially when grown upon natural media such as potato, rice and wheat grains. Spirally coiled hyphae of a somewhat similar form have been seen in *Ctenomyces* spp. and certain other Dermatophytes^(9, 14).† The natural media mentioned also provide good crops of microconidia,

* Since this paper was written definitely positive results have been obtained from inoculation experiments with the second endophyte.

† I am indebted to Dr E. J. Butler and to Mr E. W. Mason of the Imperial Mycological Institute for these references.

though these were seen sometimes on the growth arising directly from the pith lying on agar. The microconidia arise in succession from the clavate tips of closely branched, penicillate conidiophores (Pl. VIII, fig. 4). The type of branching and the method of budding off of conidia is shown in Text-figs. 12-14. The conidia, which are not



Text-figs. 10-14. The second *Lolium* endophyte in culture

10. Rope of mycelium with spiral coils.
11. Origin of a spiral; material taken from a culture on potato.
12. Microconidiophora somewhat simplified.
13. Young conidiophore formed on a hanging drop of malt agar showing the abstriction of microconidia.
14. Sterigmata and microconidia.

formed very abundantly, seldom or never remain in chains but tend to adhere together and to form with the conidiophores minute hyaline masses which become waxy-looking in dry cultures. The conidiophores usually arise from long strands of mycelium but occasionally they are formed in numbers from the cells of one of the coils (Pl. VIII, fig. 5). The conidia are hyaline, oval, and measure

$2-2.5 \times 1.5 \mu$. They resemble those associated with species of *Sclerotinia*⁽¹⁸⁾ and *Botrytis*⁽³⁾, but there has been no sign of sclerotial formation in any culture of the *Lolium* fungus.

II. THE GROWTH OF THE FIRST ENDOPHYTE IN CULTURE

All previous attempts to isolate the *Lolium* fungus have been made either with the fruit of the grass (usually *L. temulentum*) or with young seedlings; no two authors have obtained the same organism. The isolations which have been regarded as representing the endophyte are a smut⁽¹¹⁾, *Fusarium* sp.⁽⁸⁾, *Alternaria* sp.⁽¹⁾ and more recently *Chaetomium* sp.⁽¹⁰⁾.

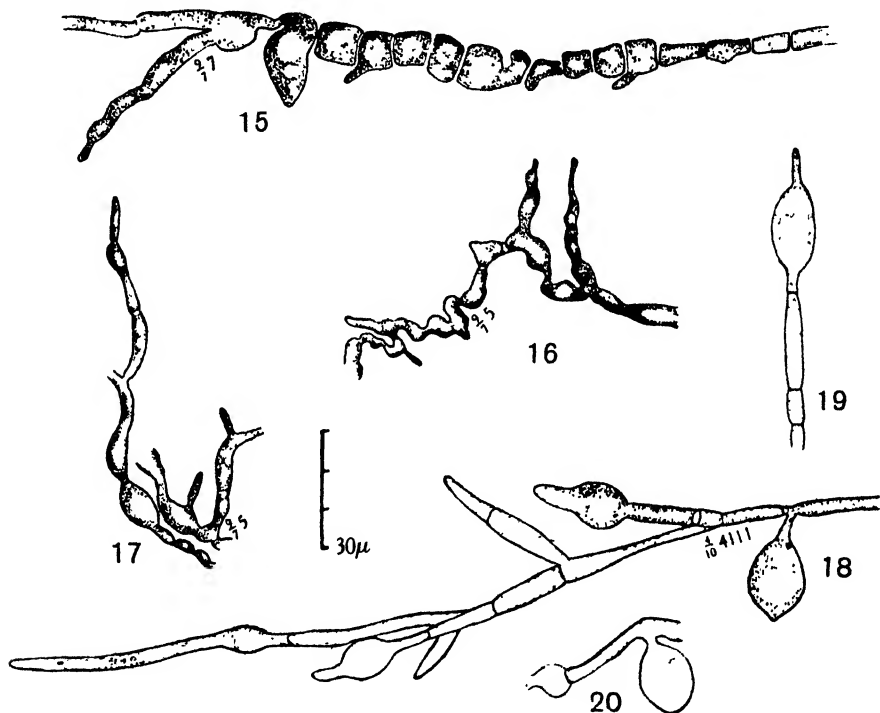
Efficient sterilization of seeds is known to be difficult. Following the technique of Guennewig⁽¹⁰⁾, grains of darnel (No. 547) carrying the endophyte were treated with a solution of bleaching powder, washed and put to germinate under aseptic conditions. Later seedlings were transferred to a nutrient solution in small flasks. Some became coated with mycelium, others remained superficially sterile and elongation of the shoot continued for some days. After a week the colcoptiles of the "sterile" seedlings were fixed and sectioned and the typical endophyte was found therein. Sterilization had clearly not affected the growth of the fungus in the plant, but it showed no signs of leaving the host tissues and growing out into the medium. This suggests that the various moulds present on some seedlings represent fungi on or within the pericarp which the sterilizing agent failed to destroy.

The presence of abundant mycelium in pith suggested that the fungus might be brought into culture by the technique which was successful with *Epichloe typhina* and the second endophyte of *Lolium perenne*. In the first trials, started in 1933, strips of pith from *L. perenne* (No. 1971) taken with every possible precaution against contamination, were placed upon the surface of an agar plate and examined at intervals under the microscope. Although the mycelium of the endophyte was clearly visible in the pith, no growth from the latter occurred. Stained preparations of strips of pith which had rested on nutrient agar for over a week showed that the hyphae had swollen and put out very short branches, but these made no further growth (Text-fig. 15). In contrast to this, pith containing *Epichloe typhina* and the second *Lolium* endophyte readily yielded their characteristic growth (p. 89).

In 1934 further tests were carried out. In collaboration with Mr C. G. C. Chesters, attempts were made to grow the fungus by placing strips of pith from infected plants of *L. perenne* and *L. temulentum* in media containing the expressed juice of darnel stems which was rendered germ-free by passage through a Seitz Werke No. 6

filter.* These experiments again gave negative results, but Mr Chesters made the valuable suggestion of trying an egg medium which Ayers found successful in cultivating *Dispira cornuta* (2).

The medium was prepared by mixing the white and yolk of a fresh egg lightly together, coagulating it by steam in a cylindrical vessel $1\frac{1}{2}$ in. in diameter, cutting slices of the "scrambled" egg column about $\frac{1}{2}$ in. thick and placing these disks in small Petri dishes. Sterilization was affected in the autoclave and the lid of each dish



Text-figs. 15-20. First endophyte of *Lolium perenne* (1971)

15. Mycelium in pith after nine days on prune agar.

16, 17. Mycelium in pith after three days on egg medium.

17. Branches with intercalary swellings.

18-20. Mycelium from sub-culture on potato-dextrose agar with intercalary swellings.

was only removed to allow one or two strips of pith to be placed on the smooth, slightly honeycombed surface of the medium.

The first experiment was started on 4 July. On the fifth day slightly opaque lines could be seen in the pith when examined under a lens. Pieces stained with cotton blue showed that growth had occurred in the long parallel strands of mycelium. Growth, which was visible to the naked eye after eight days, was at first restricted

* This section of the work was carried out in the Department of Botany of the Birmingham University in order that use might be made of the efficient press for sap extraction designed by Dr W. Leach (12). I am indebted to Prof. W. Stiles or the kind hospitality of his laboratory.

to the pith. It consisted of more or less spherical cushions which finally joined together to form a raised mass about 2 mm. in diameter, varying in length with the size of the strip of pith. Superficially the growth resembled a bacterial colony but it was firm to the touch and under a lens the surface was seen to be covered with short, erect, aerial hyphae. The growth was hyaline or cream coloured, not offering much contrast to the egg itself. In this initial experiment, each of six dishes gave identical results and none was spoilt by contaminations.

A more extensive experiment was at once set up which included malt and prune agar as well as egg medium, and pith was taken from *Lolium perenne* (No. 1971) carrying the first endophyte, *L. perenne* (No. 1972), carrying the second endophyte and infected plants of *L. temulentum* (No. 547). Pith from the first-named plant gave the same type of growth on egg but no macroscopic growth on the two kinds of agar. *L. perenne* (No. 1972) yielded cultures of the second endophyte on both egg and agar though it did not flourish on the egg medium. *L. temulentum* failed on all the media.

Further repetitions produced the same results and by staining samples of pith at intervals various stages in the growth of the mycelial strands were observed under the microscope (Pl. IX, figs. 10, 11, and Text-figs. 16, 17). Branches frequently showed the intercalary swellings which were seen in subsequent subcultures.

Transfers of the growth from *L. perenne* (No. 1971) on egg were made to fresh dishes of the same medium and some of these were left in a sterilized moist chamber during the vacation. These remained free from contamination and the cultures were multiplied in October by fresh transfers to egg.

In November an attempt was made to transfer to different media. When pieces of the growth were first placed on the surface of agar, lysis rapidly occurred in many cells but when the inoculum was rather large, some pieces of the mycelium survived and after a few weeks produced a glabrous convoluted growth on the surface of nutrient agar. When this was taken back to egg it made a colony similar to that which developed from the original pith. Transfers from egg to agar media and vice versa have been made many times with the same results. Some lysis always occurs, but most of the transfers are successful in establishing a subculture. The fungus has now been grown on a variety of media including potato disks and agar prepared with Knop's solution, Lemco, egg, liver extract, malt, potato dextrose, oatmeal and on gelatine containing egg. Growth on the last medium was sparse and little raised from the surface. Gelatine was not liquefied. On all the agar media growth was raised, forming smooth brain-like folds of about the same colour as the agar. Aerial hyphae were absent (Pl. IX, fig. 12).

Experiments in 1935 were intended to re-confirm the differential effect of egg and agar in starting initial growth, and to extend the work to other species of *Lolium*.

The first set of cultures was made on 29 May with *L. perenne* (No. 1971). Each of four dishes containing egg to which pith was added gave the same type of growth as subcultures from isolations made in 1934. No growth developed from pith placed on malt agar, and no growth was obtained on egg or malt when the pith was taken from *L. temulentum* (No. 547).

During June and July this experiment was repeated a number of times using young and old stems, and the following facts were established:

(1) The endophyte from *L. perenne* (No. 1971) was successfully isolated only when egg was used to stimulate the initial stages. Afterwards the fungus would grow on agar media provided the inoculum was sufficiently large.

(2) Negative results with pith containing abundant mycelium were always obtained with *L. temulentum*, *L. temulentum* var. *arvense* and *L. remotum*, while *L. perenne* (No. 1971) invariably yielded typical colonies on egg. There is perhaps an indication here of strain differences in the endophytic fungus of *Lolium*.

The microscopic features of the fungus in culture offer few points of interest. The growth consists of fine septate mycelia with a tendency to form rather frequent intercalary swellings (Text-figs. 18-20). Similar swellings appeared in the branches put out by mycelium in pith after three days' contact with egg (Text-fig. 17). It is unfortunate that the growth of this fungus as a saprophyte has so far not helped to establish its identity.

DISCUSSION

I. It is evident that little can be said concerning the taxonomic position of the fungus described in this paper as the second endophyte of *Lolium*. It may, however, be useful to summarize the evidence for regarding it as distinct from other known species found in association with *Lolium* and related genera of Gramineae.

Epichloe typhina which causes a closely similar type of systemic infection in *Festuca rubra* readily produces its typical conidia in cultures obtained from infected pith, and forms neither the mycelial coils nor the microconidia characteristic of the present fungus. The same holds true for *Claviceps purpurea* which was isolated from sclerotia for purposes of comparison. Mycelial characteristics in the plant and in culture, coupled with the presence of microconidia, do not suggest any relationship with the smut fungi. These features and the ease with which the fungus adopts a saprophytic existence readily

distinguish it from that which we have called the first or original *Lolium* endophyte. When these two endophytes are closely compared they seem to have no features in common except sterility and habitat.

Recent evidence that the microconidia of certain Ascomycetes may function as male gametes (4, 5, 6), and the mass of data relating to heterothallism, raise the question whether the second fungus in *Lolium* may not be a unisexual form which needs a strain of the opposite sex in order to fructify. If this be so, identification awaits the chance discovery of the right strain and no changes in the nutrient conditions of cultures now available are likely to produce the desired result. On the other hand it may be a mutant which is permanently sterile. Until new facts are discovered it seems best to use a non-committal name such as that given in the title of this paper.

The growth of the first endophyte in culture

II. The cultural work with the first *Lolium* endophyte has not thrown any new light upon the identity of this organism except in a negative sense, since it has made it rather more difficult to accept the names already suggested. The possibility remains however that closely similar types of systemic infection may be caused by quite distinct organisms.

A fresh investigation of the *Lolium* endophyte was originally undertaken in the hope of finding a link between this organism and *Epichloe typhina*, but the culture work has brought to light points of difference rather than marks of similarity and the view that they might be identical must, I think, be abandoned.

It is interesting that another organism which appeared to be an obligate parasite has been brought into culture by the use of a highly concentrated protein medium and that after living for a time as a saprophyte it has grown almost equally well on various agar media. This behaviour and the fact that it is important to transfer large pieces of inoculum coincide with the results Sawyer obtained working with some entomogenous members of the Entomophthoraceae (17).

A further point of interest is the fact that so far only the strain living on *Lolium perenne* has been brought into culture. The endophytic mycelia in *L. remotum*, *L. temulentum* and *L. temulentum arvense* have not yet been cultured away from the host. It seems likely that a more extensive search for strains of both the *Lolium* endophytes might yield interesting results.

It is, of course, very desirable that those endophytes which have been brought into culture shall be successfully reintroduced into the host. At present, belief that cultures represent the fungi inhabiting these particular grasses rests upon the direct observations of the growth of the mycelial strands in pith strips and upon the consistent

development of the same type of growth in culture. Inoculation has not yet met with success.*

SUMMARY

A description is given of the systemic infection of *Lolium perenne* by a fungus which does not fructify in the plant and which causes no pathological symptoms. It is transmitted by vegetative propagation and by seed. Mycelial characteristics, the fact that it does not form a thick zone of mycelium in the fruit and that it will grow readily as a saprophyte on all kinds of media distinguish it from the well-known endophyte of *Lolium*.

In culture, spiral coils and microconidia are formed. Since no other types of spore and no sclerotia have been seen, identification is impossible. For the present the organism is referred to as the second endophytic fungus of *L. perenne*.

Various attempts have been made to cultivate the first endophyte of *Lolium*. Using egg as a medium, cultures have been obtained which are believed to represent the fungus in *L. perenne* (No. 1971). After initial isolation on egg the fungus will grow in agar media but it remains sterile on all media tested and its culture has provided no clue to its identity.

ACKNOWLEDGEMENTS

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EXPLANATION OF PLATES VIII, IX

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PLATE VIII

Figs. 1-3. Mycelia in pith-scrapings stained with cotton blue.

Fig. 1. First endophyte from *L. perenne* (No. 1971).

Fig. 2. *Epichloe typhina* from healthy panicle of *Dactylis glomerata*.

Fig. 3. Second endophyte from *L. perenne* (No. 1972).

Figs. 4, 5. Microconidia from culture of second endophyte. In Fig. 5 the conidiophores have developed from a spiral coil.

Fig. 6. Mycelium of second endophyte lying over aleurone layer in ripe caryopsis of *L. perenne* (No. 1972).

* See footnote, p. 89.



Fig. 1



Fig. 2



Fig. 3



Fig. 4



Fig. 5



Fig. 6



Fig. 7

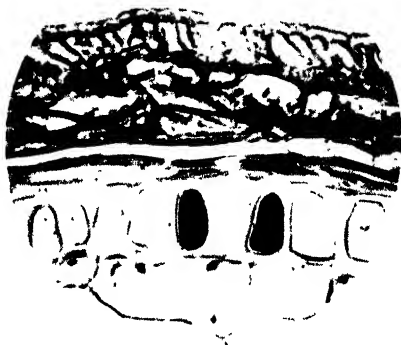


Fig. 8



Fig. 9



Fig. 10



Fig. 11

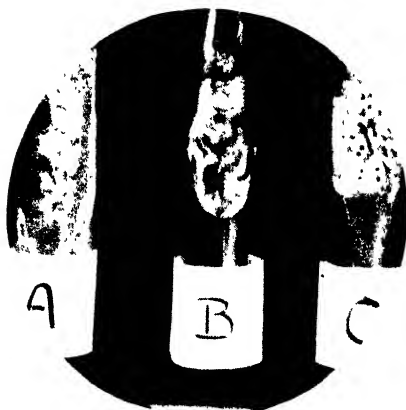


Fig. 12

PLATE IX. First *Lolium* endophyte

Figs. 7, 8. From ripe caryopses of *L. temulentum* cut transversely and stained with cotton blue.

Fig. 7. Infected female plant (No. 547) × non-infected male (No. 553).

Fig. 8. Reciprocal cross with mycelial layer absent.

Figs. 9-12. First endophyte from *L. perenne* (No. 1971).

Fig. 9. Mycelium perennating in tiller bud.

Fig. 10. Initial growth of mycelium in pith which has rested on egg for 3 days.

Fig. 11. Growth on same medium after 9 days.

Fig. 12. Growth on agar. A, potato-dextrose; B, malt; and C, ground oats.

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THE SPIRAL HYPHAE OF TRICHOPHYTON*

By A. M. DAVIDSON AND P. H. GREGORY

(With 15 Text-figures)

THE occurrence in cultures of Dermatophytes of helically coiled hyphae is a characteristic of Sabouraud's *microides* division of the genus *Trichophyton*. These spiral hyphae are not found while the fungus is living parasitically on the skin of the human or animal host, but are produced in great numbers when certain species of *Trichophyton* are grown in culture on artificial media. They are therefore organs which belong to the saprophytic phase of the Dermatophytes.

Matruchot & Dassonville (1899) drew attention to the resemblance between the spirals of the dermatophytes and the spiral ornaments, so-called, which surround the perithecia in *Ctenomyces serratus* Eidam. This resemblance was used by Langeron & Milochevitch (1930) to justify the inclusion of the Dermatophytes in the family Gymnoascaceae, although asci or functional perithecia are usually absent, and not known with certainty.

Little attention has been paid to these remarkable structures, though Grigoraki (1927) made their presence the basis of his genus *Spiralia*, in which he separated off the members of Sabouraud's group *microides*. The validity of this distinction was shaken when Langeron & Milochevitch (1930) showed that on media such as grains of oats *Microsporum felineum* could be induced to produce spirals which were up to that time unknown in that species. Emmons (1934) occasionally found spirals in cultures of the same species even on Sabouraud's maltose-peptone agar (*milieu d'épreuve*).

Spirals have been regarded usually as ornaments or curiosities of no particular significance. Thus Emmons says "like some other structures produced by the fungi, they are not believed to possess any particular function or evolutionary significance". The investigation here described was prompted by the idea that organs formed in such regularity and abundance as the spirals of *Trichophyton gypsum* are unlikely to be produced by such plastic organisms as the fungi without some significance in the life history. The long supposed functionless pycniospores of *Puccinia* may be cited as an example of the fallacy of regarding constantly occurring fungus organs as rudimentary. The

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development and structure of the spirals were studied, and as a result of experiments on the properties of mature spirals a tentative suggestion as to their function is put forward.

Although no Dermatophyte has yet been observed growing as a natural saprophyte, evidence is accumulating to indicate that under suitable conditions they can enter upon a saprophytic phase of growth. Recent investigations along this line have been reviewed by Gregory (1935). At present the saprophytic phase is known only under laboratory conditions. A study of spiral structure may therefore be expected to throw light on the nature of the saprophytic phase of these fungi.

DEVELOPMENT OF SPIRALS

The stages in the development of spirals on a colony may be illustrated by the following description of a culture on wheat-flour agar. The culture was a typical strain of *Trichophyton gypseum* which had been derived from a single conidium (aleuriospore) isolated by the wet needle method of Hanna (1928). As noted by previous writers, these monoconidial cultures developed conidia (aleuriospores), macroconidia (fuseaux), and spirals as freely as cultures derived from mass inocula (Spring, 1931; Emmons, 1931). The medium was prepared by adding agar to a paste containing 1 per cent whole-wheat flour in water, and heating on a water-bath for several hours to remove air before sterilizing in conical flasks in the autoclave. Removal of the air was necessary to prevent frothing of the medium in the small containers.

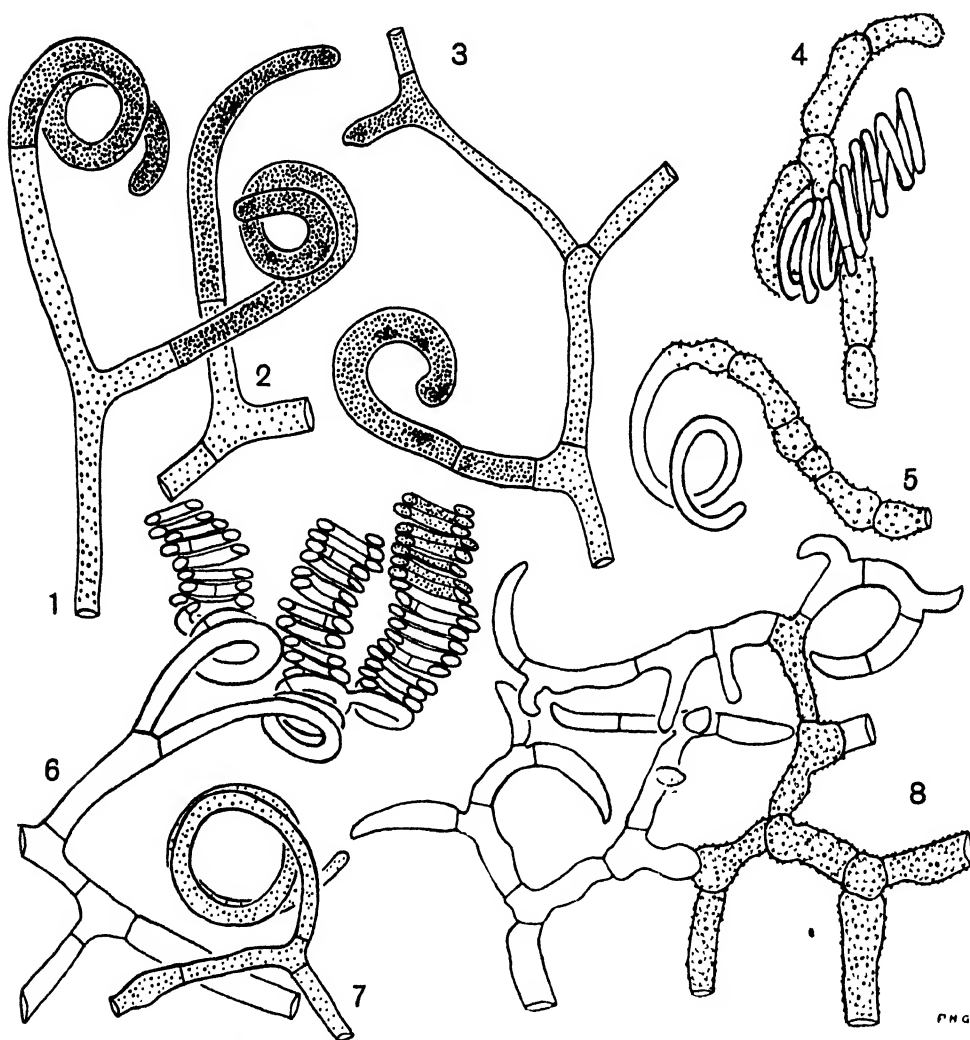
The cultures were kept at room temperature for four weeks, and at the end of that time had a diameter of 6 cm., corresponding to a radial growth of 7-8 mm. per week. The cultures had a chalky granular surface which showed pronounced zonation. From each zone portions of mycelium were removed and examined microscopically. The occurrence of spirals in the various zones is shown in Table I.

Table I. *Occurrence of spirals in various zones of culture of Trichophyton gypseum*

Zones	Condition of spirals
(1) Outer translucent zone of radially growing pioneering hyphae immersed in medium, forming edge of colony	No spirals
(2) Elevated white downy zone free from spores	No spirals
(3) White powdery zone about 1 cm. in breadth containing numerous conidia (aleuriospores)	Spirals developing as in Figs. 1-3
(4) Cream-coloured powdery zone occupying greater part of colony	Mature (empty) spirals abundant
(5) Downy central tuft over inoculum	No spirals formed

From examination of different parts of the third zone it was concluded that conidia are produced somewhat earlier than spirals, since

conidia were found nearer the edge of the colony than were spiral initials. The zone of active spiral production was estimated to be from seven to ten days old.



Figs. 1-6 and 8. Development and structure of spirals of *Trichophyton gypseum*.

Figs. 1-3. Stages in development of spiral and establishment of asymmetry. Culture on 1 per cent wheat agar. ($\times 1100$.)

Figs. 4, 5. Mature spirals borne on torulose hyphae. Culture on wheat agar. ($\times 485$.)

Fig. 6. Well-developed branched spirals from *in situ* culture on hair naturally infected by *T. gypseum*, kept in a saturated atmosphere. The distal portions of the spirals represented in section to show flattening of component hypha in a plane at right angles to the spiral axis. ($\times 1100$.)

Fig. 7. An aberrant spiral of *T. interdigitale* with clockwise asymmetry. ($\times 485$.)

Fig. 8. Crooked hyphae often associated with spirals on mycelium. ($\times 1100$.)

The spirals begin either as short lateral branches or as terminal cells on aerial hyphae which have somewhat denser protoplasm than the ordinary hyphae. Most commonly the spirals arise in a lateral

position on an aerial hypha. One of the cells of the hypha develops a growing region, and puts out a lateral diverticulum at right angles to itself; this arises somewhere between the middle of the cell and the distal septum which separates the cell from the younger cell in front. A lateral hyphal tip is thus established and growth continues. The new hypha is not as a rule cut off from the parent cell by a septum at its base, but by a septum laid down a short distance along the hypha. The basal cell of the branch thus becomes T-shaped. When the second or third septum has been formed the tip of the hypha begins to curve to one side, and this curvature is continued in all successive cells of the spiral. Continued growth around an arc would soon bring the tip of the hypha into contact with the older portion of the same hypha. However, before this occurs a lateral displacement of the growing tip becomes evident, and the tip passes, in the course of its growth, to one side of the previously formed part of the curved hypha. An asymmetrical structure is thus produced (Fig. 1). As will be shown later, it is evident from observations on mature spirals that this lateral displacement of the growing tip is always to one particular side of the older hypha.

The developing spiral becomes cut off from the parent hypha by a septum, and as the spiral grows at the tip its protoplasm moves forward, leaving the hypha empty behind. Stages were found in fixed preparations in which the older parts of the structure were empty, only the cell walls being visible, while the growing tip was full of refractive protoplasm which stained with methylene blue (Fig. 6). Direct observations on living spiral hyphae showed that they made from a half to one complete turn in twenty-four hours.

STRUCTURE OF MATURE SPIRALS

Fully mature spirals are empty tubes; they fail to take up cytoplasmic stains. Presumably the protoplasm has been exhausted in the laying down of the spirally coiled wall. In one experiment two mature spirals were removed from the mycelium with needles, and placed on Sabouraud's agar. They were observed at intervals under the microscope for several days, but as they showed no signs of germination, they are regarded as dead structures. The emptying of the older part of the hypha is common in the Dermatophytes as in many other fungi. Sabouraud (1910) noted that the protoplasm in sporogenous hyphae became encysted in the spores, leaving the conidiophore empty. There is a striking difference between an emptied spiral and an emptied conidiophore. A conidiophore, or an ordinary vegetative hypha, when emptied of its protoplasm is a shrunken irregular structure which tends to collapse. When mounted in a refractive medium such as glycerine jelly it is almost invisible. The emptied portion of the spiral,

on the other hand, is a smooth, strong, easily visible tube, showing no tendency to collapse or become irregular in outline. The wall of the spiral is apparently thicker and more durable than that of the conidiophore. Spiral hyphae have not so far been observed to bear spores, though they usually arise in close proximity to spore-bearing hyphae. They are always aerial. The hypha which bears the spiral may be stout and torulose (Fig. 4). Sometimes in place of spirals these torulose hyphae bear peculiar crooked hyphae (Fig. 8) of unknown significance.

When mature and empty, the spirally coiled hypha is flattened from side to side. The cross-section of the component hypha is an ellipse whose major axis is at right angles to the axis of the spiral. This flattening can be seen by focusing upon a mature spiral which may be either in air or mounted in glycerine jelly. When the objective is focused on the plane of the axis of the spiral so that the component hypha is seen in optical section, it is apparent that the wall of the hypha is flattened above and below as shown in Fig. 6. Whether the flattening arises by mutual pressure of the coils of the spiral during growth or as a result of unequal growth of the cell wall is not known. The flattening probably has a mechanical significance which will be dealt with when the possible function of the spirals is considered below.

In the older parts of a culture, spirals of from one to thirty turns can be found. The greater number of turns has been found in specimens prepared by the *in situ* culture technique (Davidson & Gregory, 1934).

When the mycelia are growing under favourable conditions some of the spirals may be branched, a new growing tip being established at some point on the spiral curve. Growth is at first radially away from the parent spiral, but soon the tip of the hypha bends round and forms a second spiral alongside the first. This process may occur several times, especially in cultures prepared by the *in situ* technique (Fig. 6). The occurrence of branched spirals does not appear to have been observed by previous writers.

By counting the number of coils (n) and measuring the inside diameter (d) of a spiral at right angles to the spiral axis, it is easy to calculate approximately the length of the hypha composing the spiral. When the successive coils of the spiral are in contact it may be assumed that each coil is a circle, and that the length (L) of the component hypha will be approximately given by the formula: $L = n\pi d$. d is taken as the inside diameter, as it is the inside or shorter side of the coil which will limit the extension of the hypha. In many spirals the successive coils are separated instead of being in contact; the length of the hypha as calculated from the formula will then be too low. The dimensions of seven hyphae which were selected at random

from three cultures are given in Table II, from which it will be seen that the hypha may be up to thirty times as long when extended as when coiled up in the spiral.

Table II. *The length of the hypha composing the spiral*

Isolate No.	Length of spiral μ	No. of coils	Inside diameter μ	Length of hypha (calculated) μ	$\frac{\text{Length hypha}}{\text{Length spiral}}$
306	13	6	11	250	19 : 1
306	26	15	15	760	29 : 1
306	39	10	15	500	13 : 1
306	16	7	10	240	16 : 1
306	33	12	12	500	15 : 1
377	21	14	8	350	17 : 1
549	48	28	8	700	15 : 1

The number of spirals developed upon a mycelium is very great. On a medium prepared by diluting Sabouraud's maltose-peptone agar with nine times its volume of plain agar the spirals were relatively sparse and could be counted. A square millimetre of the surface of a mature colony on this medium was estimated, by counting the spirals visible on a small area under a low-power objective, to bear about five hundred spirals. In various isolations and on other media considerably different figures are likely to be found, but the number of spirals produced is certainly great.

ASYMMETRY OF THE SPIRALS

In the description of the development of a spiral given above it was noted that as the tip of the hypha grows along the curved path which constitutes the first turn of the spiral, the tip becomes displaced to one side so that, instead of meeting the previously formed part of the hypha, it passes to one side of the older part of the hypha. An asymmetrical structure is thus produced, and all further growth of the hypha perpetuates this asymmetry. Further, it was noticed that, with very rare exceptions, the growing tip always passes to one particular side of the base of the spiral, so that the asymmetry of the resulting spiral is counter-clockwise.* That this counter-clockwise structure is not merely an attribute of one culture is shown by the list in Table III of isolations derived from various sources, in all of which counter-clockwise coiling was the rule.

* The sense in which the term "counter-clockwise" is here used is defined below in a section on the terminology of spiral structure. The meaning may be illustrated by saying that counter-clockwise spirals of *T. gypsum* resemble the counter-clockwise spirals of the scarlet-runner or *Convolvulus*, but differ from the clockwise spirals of the hop and honeysuckle among the twining plants.

Table III. *Cultures bearing spirals with counter-clockwise asymmetry*

Species	Source
<i>Trichophyton gypseum</i>	(144) <i>Sycosis parasitaria</i> , Manitoba (306) <i>Tinea corporis</i> , Manitoba (376) <i>Kerion Celsi</i> , Manitoba (377) <i>Kerion Celsi</i> , Manitoba (549) <i>Tinea corporis</i> , Manitoba
<i>Trichophyton asteroides</i>	American Collection of Type Cultures
<i>Trichophyton granulosum</i>	American Collection of Type Cultures
<i>Trichophyton (Epidermophyton) interdigitale</i>	Dr E. Muskatblit, New York (250) <i>Tinea pedis</i> , Manitoba (564) <i>Tinea pedis</i> , Manitoba (615) <i>Tinea pedis</i> , Manitoba (624) <i>Tinea pedis</i> , Manitoba (635) <i>Tinea pedis</i> , Manitoba
<i>Trichophyton persicolor</i>	National Collection of Type Cultures, London

The counter-clockwise structure was first observed in cultures on Sabouraud's maltose-peptone agar. To see whether nutrient substances influenced the direction of coiling, several strains were tested on a variety of other media, but no alteration was observed in the direction of twist. The media used were: (1) Sabouraud's maltose-peptone agar diluted with nine times its volume of plain agar; (2) Difco "Sabouraud's dextrose agar"; (3) Sabouraud's maltose-peptone agar; (4) wheat-flour extract agar; (5) whole-wheat flour paste; (6) human hair in moist tubes; (7) *in situ* cultures on naturally infected human hair; (8) sterilized horse dung; and (9) sterilized cow dung. On Czapek's solution-agar growth was poor and spirals were not produced.

External factors did not appear to affect the structure. Counter-clockwise spirals developed as normally in cultures grown in total darkness from the time of inoculation, as in those which were exposed to the usual diurnal alternation of light and darkness in the laboratory. The spirals also appeared to be insensitive to gravity, as the counter-clockwise rule held whether the spiral had grown vertically upwards, downwards, horizontally, or in intermediate positions.

The effect of temperature cannot be dealt with in the same way. When strains of *T. gypseum* and *T. interdigitale* isolated in Manitoba were grown on Sabouraud's maltose-peptone agar in the incubator at 37° C. spirals were not observed. This suppression of the formation of spirals at body temperature perhaps gives a clue to one of the factors responsible for the differences in the morphology of *T. gypseum* in its saprophytic and parasitic phases.

All the isolates examined so far have shown counter-clockwise spiral structure. Exceptions have been in the occasional occurrence of a few clockwise spirals in certain cultures in which, however, the majority of spirals were counter-clockwise. These clockwise spirals

are rare, and were observed in only two cultures of *T. interdigitale* isolated in Manitoba. In one of these cultures, out of 137 spirals carefully examined with the oil-immersion lens, only eight (less than 6 per cent) showed clockwise structure. In the other culture the proportion observed was only 3 per cent. It is possible that all cultures tend occasionally to produce clockwise spirals. It should be remarked that the clockwise spirals were well formed and normal in appearance. Apparently their production depended not so much on the breaking down of the normal mechanism for producing counter-clockwise spirals, as on a complete reversal of that mechanism. Perhaps this phenomenon will one day be explained as due to the local occurrence in the mycelium of a stereo-isomer of a normal molecule.

EXPERIMENTS ON THE PROPERTIES OF THE SPIRALS

The properties of spirals were studied experimentally by means of a simple micromanipulator consisting of a glass needle or hair attached at an angle of about 30° to a horizontal glass rod. The glass rod was mounted in a cork in place of the objective of a microscope. The focusing device of this microscope provided a delicate vertical movement of the needle. A Petri-dish culture of *T. gypsum* was placed for observation on the stage of another microscope. By use of the mechanical stage of the observation microscope, motion in two other directions was possible. By means of this micromanipulator the tip of the glass needle or hair could be made to touch any part of the mycelium in the Petri dish. This simple micromanipulator was suggested to one of us in 1930 by Dr T. R. Vernon.

In one experiment the properties of spirals were studied by touching them with glass needles of $30\text{--}40\mu$ diameter which had been cleaned in alcohol and ether in order to free them from grease. If ordinary straight vegetative hyphae were touched with a needle, hypha and needle adhered. When by means of the micromanipulator the needle was withdrawn the adhesive force was sufficient to deform the needle. However, on withdrawing the needle still further the adhesion was broken and both hypha and needle swung back to their former positions. Adhesion between hypha and needle was insufficient to break the hypha free from the mycelium.

If the needle were made to touch either the tip of a spiral or the middle of a coil the same phenomenon was observed, but owing to the uncoiling of the spiral the needle had to be withdrawn further before the hypha was fully extended and adhesion broken. The spiral hypha when drawn out was seen to be flattened and appeared as a chain of arcs. This form would be expected from the flattening of the hypha already noted above. When adhesion between hypha and needle had been broken the hypha usually regained its spiral form completely.

The peculiar flattening of the spiral hypha in a plane at right angles to the axis of the spiral no doubt has a function in restoring the hypha to the original shape after it has been temporarily deformed. On account of this flattening there will be more resistance to any deformation of the curvature of the hypha, and less resistance to distortion of the coil in the direction of the axis of the spiral.

If the needle was made to touch a number of coils of a spiral, or to touch several adjacent spirals at the same time, events followed a different course. On withdrawing the needle adhesion was sufficient to detach a group of spirals, together with conidiophores and conidia (aleuriospores), from the mycelium, and to leave them adherent to the needle.

In place of glass needles human hairs were used in repeating these experiments, and similar results were obtained. The general effect of the spirals may be described as presenting a large adhesive surface to foreign bodies and so tending to break off a portion of the mycelium.

The micromanipulator method was also applied to cultures of *Microsporum felineum*, on whose mycelium as a rule no spirals are present. The hyphae in this species adhered to the needle only feebly, and repeated attempts failed to detach a single hypha or fuseau by adhesion to the needle.

The use of microscope and micromanipulator was not necessary in order to demonstrate the adhesive properties of the mycelium of *Trichophyton gypseum*. If an ordinary steel needle be touched lightly to the granular surface of a colony growing on agar and then withdrawn it can be observed with the naked eye that a tuft of white hyphae and spores about a millimetre in diameter has adhered to the tip of the needle. On the other hand, a needle touched to the surface of a culture of *Microsporum felineum* does not bring away a tuft of hyphae but only serves to compress together the hyphae of the culture. It is therefore obvious that a mycelium of *Trichophyton gypseum* has adhesive properties which are not possessed by a mycelium of *Microsporum felineum*. That this adhesiveness in *Trichophyton gypseum* is not due solely to the presence of spirals is shown by the fact that a "pleomorphic" culture of the species which had ceased to produce spirals still exhibited some adhesiveness when touched with a needle. The vegetative hyphae of *T. gypseum* appear to be sticky. There is nothing at present known to suggest that the spirals are more sticky than the vegetative hyphae, but the presence of spirals on the mycelium must ensure that any foreign body such as an animal touching the mycelium comes into contact with a very large area of sticky surface and leaves with a relatively large inoculum consisting of spirals, hyphae and spores firmly anchored. The spirals appear to function as local concentrations of a large quantity of sticky mycelium.

The tarsi of house flies which were induced to walk over cultures

of *T. gypseum* at once became coated with white masses of hyphae. The flies were then made to walk across the surface of sterile agar slants, and on incubating these, cultures of *T. gypseum*, recognizable by the characteristic spirals and other organs, were obtained.

There seems no reason to believe that the spirals of *Trichophyton* are functionless; rather they must be looked upon as structures serving to attach a large number of the spores which are produced while the organism is growing in its saprophytic phase to any animal which happens to come into contact with the mycelium.

DISCUSSION

The fact that a clockwise or a counter-clockwise asymmetry is usually specific for twining plants among the Phanerogams has long been familiar to botanists. Spirally twisted organs of various kinds also occur in certain fungi, but in this group, where microscopic examination is necessary to elucidate the spatial structure of the spiral, less attention has been paid to the subject. Spiral structure in the genus *Trichophyton* is worthy of study, since differences in such a clear-cut character, if found between distinct species, might be of value for the easy identification of members of this difficult genus. A method of determining the spatial structure of microscopic objects by differential focusing has been worked out and is presented as an Appendix to this paper in the hope that observations will be made on Dermatophytes and other Thallophyta by other workers. In this connexion it will be necessary to discuss the confused nomenclature of spiral structure, and to define the terms used, since studies of spatial structure are misleading if expressed in ambiguous terms.

Nomenclature

The plane spiral. For a plane spiral, whose coils lie in one plane as illustrated in Fig. 9, there is only one possible spatial structure. The apparent existence of two forms is due merely to the position of the observer, as may be shown by examining Fig. 9 through the back of the page against the light. Among fungi spores in the form of a plane spiral occur in certain members of the Helicosporeae.

Three-dimensional spiral or helix. For a three-dimensional spiral there are two possible spatial structures which differ absolutely irrespective of the position of the observer. The nomenclature of these forms is confusing, as workers in different fields of science, or even in the same field, have used terms in opposite senses. This confusion is discussed by Jackson (1916).

For anatomical reasons the head of a screw which is being driven into a piece of wood can be rotated by the right hand with more force in a clockwise direction than in a counter-clockwise direction. The

common wood-screw is accordingly threaded so that when twisted in a clockwise direction by the right hand it will be driven forward away from the operator. Such a thread, which is illustrated in Fig. 10, is called by the engineer a right-handed thread, and this logical terminology appears to be followed consistently by zoologists and human anatomists, and also by certain botanists. Unfortunately many botanists, including Darwin, Hooker and Strasburger, applied the term right-handed to twining plants whose spatial structure corresponds with the engineers' left-handed thread. The confusion which already exists in the nomenclature is illustrated in Table IV.

Table IV. *Differences in nomenclature of spiral structure*

Engineers, zoologists, anatomists and earlier botanists		Many later botanists	
Right-handed	}	=	Left-handed
Dextrorse			Sinistrorse
Dextral			Counter-clockwise
			Anti-clockwise
			Against the sun*
Left-handed	}	=	Right-handed
Sinistrorse			Dextrorse
Sinistral			Clockwise
			With the sun*

* These terms are ambiguous as they have opposite meanings in Northern and Southern Hemispheres.

Inspection of Table IV shows that if a structure is to be described as right-handed or left-handed it is essential to state clearly what system of nomenclature is being followed. In the present work the terms *clockwise* and *counter-clockwise* have been adopted as being less likely to lead to confusion than *right-handed* and *left-handed*. The definition of clockwise and counter-clockwise structure given in the following paragraph has been based on the concept of an observer viewing the spiral as a static structure. In order to make the definition of general applicability all ideas of growth or movement of the spiral have been avoided.

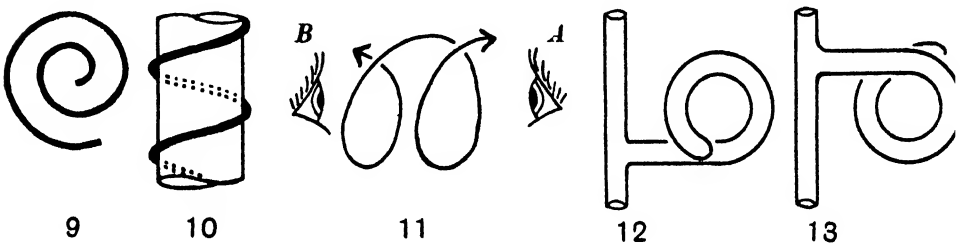
Definition. An observer situated outside the spiral, and on its axis, describes the structure as *clockwise* if the end of the coil nearer him points in the direction of rotation of the hands of a clock. Conversely, when the end of the coil nearer him points in a direction contrary to rotation of the hands of a clock the spiral is said to be counter-clockwise.

Fig. 11 illustrates a spiral of which the observer situated at *A* regards the end of the coil nearer him as pointing in the clockwise direction. The end of the coil farther from him, at *B*, will appear to him to point in the opposite direction (counter-clockwise). It is therefore important that the convention should be applied only to the

direction in which the end of the thread nearer the observer points. However, a second observer situated at *B* will regard the end of the coil at *B* as clockwise, and that at *A* as counter-clockwise, simply because he is using his own clock as reference and not that of the observer at *A* seen in reverse. Therefore by following the convention an observer situated at either end of the spiral will reach the same conclusion with regard to its spatial structure.

Spiral organs in the fungi

Spiral organs occur in a variety of fungi. Both clockwise and counter-clockwise forms occur, but, so far as attention has been paid to the subject, the evidence indicates that for any particular species the spatial structure is constant. Whether structural similarity ex-



Figs. 9-13.

tends to the wider groups it is at present impossible to say. The following examples indicate the constancy of spatial structure within the species.

Drechsler (1919) observed that rotation in the formation of spirally coiled sporogenous hyphae in the genus *Actinomyces* is specifically "sinistrorse" [counter-clockwise] or "dextrorse" [clockwise]. Out of sixteen species in which the condition could be ascertained with certainty, eleven were counter-clockwise, and five were clockwise. Drechsler drew attention to the absolute constancy with which a species adheres to one kind of rotation.

In the Myxomycete genera *Trichia* and *Hemitrichia* the capillitium threads in the sporangium have one or more helically coiled bands of thickening on the cell wall. These show a consistently clockwise structure and are so figured by Lister (1925). It is, however, interesting to note that an aberrant specimen of *Hemitrichia leiotrichia* is recorded by Miss Lister with dextral [counter-clockwise] spirals.

In Zopf's (1881) monograph of the genus *Chaetomium* spirally coiled hairs are depicted around the ostiole of the perithecium in several species. Definite statements as to their spatial condition are lacking, except that in *C. crispatum* Fuckel the coils are said to alternate in direction along each hair. The species must be almost unique in this respect. Zopf's carefully drawn figure of *C. spirale* clearly shows a

preponderance of counter-clockwise spirals. A culture of *C. ? spirale* which had been isolated from Manitoban soil and supplied by Dr G. R. Bisby when examined by us showed only counter-clockwise perithecial hairs.

A study of the figures in Linder's (1929) monograph of fungi belonging to the Helicosporeae indicates that the condition differs between different species. A specimen of *Helicoon ellipticum* (Pk.) Morgan in the Manitoba Agricultural College Herbarium was found by us to possess counter-clockwise spores.

These observations indicate that any particular collection or isolate of a fungus shows a high degree of uniformity in the spatial structure of its spiral organs. Further, this uniformity commonly extends to other individuals of the same species. It appears probable that spiral spatial structure is specific except for occasional aberrations. There is need for more data on this subject, and it is essential that the data should be expressed in unambiguous terms.

The spiral hyphae of *Trichophyton* agree with similar organs observed in other fungi in showing a high degree of uniformity of spatial structure. Further, in the possession of predominantly counter-clockwise spirals they resemble *Chaetomium spirale*, *Helicoon ellipticum* and certain *Actinomyces*, but differ from the Myxomycetes *Trichia* and *Hemitrichia*.

SUMMARY

The spiral hyphae of *Trichophyton* are believed to be organs of attachment playing a part in the dissemination of the fungus and in infecting animal hosts from saprophytic sources. Mature spirals are empty, dead hyphae, flattened in a plane at right angles to the axis of the spiral. They may be branched.

The spirals observed in five isolates of *T. gypseum*, six of *T. interdigitale* and one each of *T. asteroides*, *T. granulorum* and *T. persicolor* were found to be counter-clockwise. No isolates were found with clockwise spirals predominating. Occasional clockwise spirals were however found in some isolates of *T. interdigitale*.

When deformed by mechanical contact a spiral regains its original shape, and it is probable that this process is aided by the flattening of the spiral hypha.

The mycelium of *T. gypseum* is sticky and tends to adhere to objects which come into contact with it. This stickiness is not shared by *Microsporium felineum*. When a sufficient number of adjacent hyphae or coils of a spiral adhere to an object they can break away from the mycelium. It is concluded that the function of the spirals is to present a large adhesive surface to any foreign body. Thus a relatively large inoculum is attached to an animal which touches the saprophytic mycelium.

APPENDIX

The determination of spiral structure by differential focusing

The method of determining the spiral structure of microscopic objects by differential focusing consists in observing the object under the microscope at such a magnification that only a portion of the spiral appears in sharp focus at one time. Then by differential focusing the spatial relations of the parts can be made out.

Two principal positions are encountered in examining spiral structures under the microscope: (1) when the observer is situated on the axis of the spiral; and (2) when the observer is situated at right angles to the axis of the spiral.

(1) *Observer situated on the axis of the spiral.* By focusing on the nearer end of the coil the structure of the spiral can be read off at once by following the definition given on p. 108. When the object is a growing structure attached at one end, such as a fungal hypha, two conditions arise according as the structure has grown towards or away from the observer.

When the spiral has grown vertically upwards towards the observer no difficulty will arise in applying the definition (Fig. 12). On focusing downwards the end of the coil nearer the observer first comes into focus, and from this the structure of the spiral can be determined at once. On focusing successively lower planes the further end of the coil is reached, and is found to point in the direction opposite to the nearer end.

When the spiral has grown vertically downwards away from the observer, the attached end of the coil which is nearer the observer comes into focus first on focusing downwards (Fig. 13). According to the definition the direction in which this end points is taken as giving the structure of the spiral; usually this direction will be opposite to that in which growth has taken place during the formation of the structure; this fact is, however, irrelevant according to the definition. On focusing down farther the tip of the spiral comes into focus and will be found to point in a direction contrary to that from which the structure is named. An observation of the more distant end of the spiral can sometimes be made use of when a good focus cannot be obtained on the nearer end.

(2) *Observer situated at right angles to the axis of the spiral.* When the observer looks at the spiral radially or "sideways-on", it is not necessary to find the ends of the coil in order to determine the structure. To an observer situated radially with respect to the axis the coils will appear to cross the axis at some angle other than a right angle. Further, if the coils lying between the observer and the axis cross the axis at an angle θ° on the right side of the axis, then the coils lying

below the axis will make an angle θ° on the left side of the axis as shown in Fig. 14, where a portion of a counter-clockwise spiral is illustrated. On focusing down slowly on to such a spiral the portions of the coil *ab* and *cd* in the plane nearer the observer come into focus first; on focusing below the axis the portion *bc* comes into focus. By considering the axis as running north and south we can express the structure by saying that in a counter-clockwise spiral the coils nearer the observer run north-east and south-west relative to the axis, while on focusing down it can be seen that the farther parts of the coil run north-west and south-east. Conversely, in a clockwise spiral the coils above the axis run north-west and south-east, while those below the

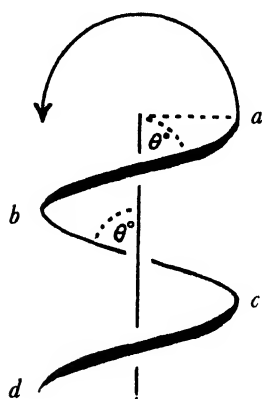


Fig. 14.

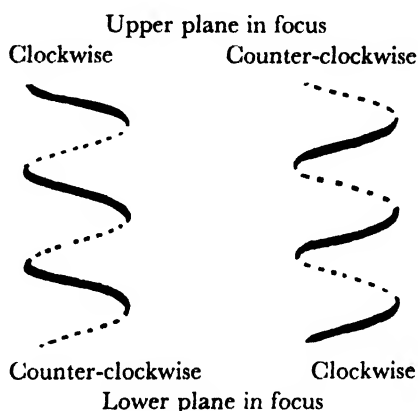


Fig. 15.

axis run north-east and south-west. The relations between the nearer and farther planes of focus in both clockwise and counter-clockwise spirals is compared diagrammatically in Fig. 15, in which those parts of the spiral which are in focus in each plane are represented by thickened lines. The compound microscope rotates the image through 180° in the plane of the field, and consequently the microscope gives a true picture of spiral spatial structure. By the method of differential focusing here described the spatial structure of a spiral can be determined by observations made on a very small portion of the spiral examined in any position.

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SOME OBSERVATIONS ON THE OCCURRENCE OF *FUSARIUM CULMORUM* ON WHEAT

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IN published work dealing with the occurrence of *Fusarium culmorum* on cereals most attention has been devoted to the status of this fungus as a parasite. Simmonds(7), Bennett(1), Russell(5), Geach(3) and others have recorded it as causing seedling blight, foot rot, whiteheads or ear blight of oats and wheat in the field, and have proved its pathogenicity by controlled inoculation experiments. In recent years, however, references to its occurrence either as a weak secondary parasite, or under circumstances where its parasitic action has been doubtful, have been more numerous. One of the most interesting of these is the work of Broadfoot(2) in which isolations from many thousands of wheat plants taken at random from rotation plots yielded *F. culmorum* (either alone or in combination with other fungi and bacteria) to the extent of from 20 to 60 per cent in different plots. Simmonds(7) and Sanford & Broadfoot(6) have also isolated *F. culmorum* from stubble. The latter workers compared the virulence of many isolates and found that most of them were only weakly pathogenic. Apart from this recent work in Canada the information available on the weakly parasitic and saprophytic life of *F. culmorum* on cereals is comparatively scanty. It therefore seems worth while putting on record the results of isolations made from healthy wheat plants collected in England.

Isolations made at Rothamsted about harvest time in 1934 showed that *F. culmorum* was present on most of the roots and stem bases of wheat plants affected with whiteheads. However, control isolations made from healthy plants with full ears of grain revealed that *F. culmorum* was often present on the roots and stem bases of these plants also, so that no conclusion could be drawn as to the parasitic action of the fungus where it had been isolated from the whitehead plants.

In the following year, with a view to obtaining some idea of the progressive invasion of the roots and crowns of wheat plants by fungi, isolation work was begun about the time the crop was in flower, and periodic isolations were made as the crop ripened. The material for this study was obtained from three wheat fields, one at Rothamsted, one at St Albans and one at Cambridge (Table I).

The crops on these fields were among the healthiest in the districts,

Table I. *Cropping data for fields examined*

	Soil	Variety	Sown	Reaped	Rotation	Yield per acre
Rothamsted	Clay with flints, pH 7.8	Victor	26 Oct. 1934	6 Aug. 1935	1932 kale 1933 barley 1934 beans	
St Albans	Clay with flints, pH 7.2	Victor	27 Sept. 1934		Temporary grass ley 1934 oats	50 bushels
Cambridge	Heavy gault clay, pH 7.9	Yeoman I	29 Oct. 1934	30 Jul. 1935	Temporary grass ley	58 bushels

and showed no evidence that seedling diseases had been present. At intervals of about a fortnight ten samples of about a dozen plants were taken from each crop. These samples were taken at intervals of ten paces and between ten and twenty paces in from the edge along the best side of the field. From each sample the healthiest plant was chosen, and four crown pieces and five root pieces cut out at random. These were rinsed in alcohol, sterilized two minutes in 1:1000 mercuric chloride, washed in sterile water, and plated out on potato-dextrose agar, the crown pieces and root pieces being plated in separate dishes. The last isolations were made from the stubbles after the crops had been cut. A summary of the results is given in Table II.

Table II. *Isolations of Fusarium culmorum and other fungi from healthy wheat plants*

Field	Date of collection	Part of plant	No. of pieces	Number of pieces yielding		
				<i>Fusarium culmorum</i>	other <i>Fusarium</i> spp.	other fungi and bacteria
Rothamsted	26 June	Crown	40	0	4	29
		Roots	50	5	2	36
	7 July	Crown	40	4	1	31
		Roots	50	2	2	41
	22 July	Crown	40	3	5	32
		Roots	50	2	3	45
	6 Aug. (stubble)	Crown	40	1	0	35
		Roots	50	0	1	46
St Albans	1 July	Crown	40	0	0	17
		Roots	50	1	0	15
	15 July	Crown	40	1	5	33
		Roots	50	5	3	36
	29 July	Crown	40	14	3	21
		Roots	50	10	2	33
	13 Aug. (stubble)	Crown	120	42	2	66
		Roots	150	34	0	106
Cambridge	12 July	Crown	40	13	0	23
		Roots	44	14	0	21
	26 July	Crown	40	11	4	25
		Roots	150	14	5	75
	20 Aug. (stubble)	Crown	120	90	0	4
		Roots	150	130	0	6

It will be seen that *F. culmorum* was isolated from a few specimens in every field at the time of the first sampling. In the Rothamsted field it did not increase in amount as the season advanced; in fact, it decreased. In the St Albans and Cambridge fields, on the other hand, it increased markedly, and in each, seven of the ten plants taken before harvest had the fungus on stem base or roots. In the field at St Albans thirty stubble samples taken at random over the field two weeks after harvest showed 70 per cent infection with *F. culmorum*, and in the Cambridge field the fungus was present on every one of fifty random stubble samples taken three weeks after harvest.

In spite of the appreciable amount of *F. culmorum* present on the roots at harvest time the crops gave excellent yields. The one at St Albans yielded 50 bushels to the acre, and the one at Cambridge 58 bushels. In the latter there were fourteen sacks of head corn and only two-thirds of a sack of tail corn, so that the grain had filled remarkably well. A sample of wheat from this crop took the Gold Cup at the Baker's Exhibition, London, for the best sample of milling flour.

To determine whether the strains of *F. culmorum* obtained were weak and almost non-pathogenic, isolates from the different localities were multiplied on corn-meal sand, and pathogenicity tests on wheat seedlings in the greenhouse were carried out in the usual manner. When the seedlings were washed out at the end of five weeks there was appreciable attack on the roots by all the strains of *F. culmorum* used. The average disease rating (4) for thirteen isolates of *F. culmorum* was 21.0 (ranging from 9 to 40), that for three isolates of *Ophiobolus graminis* was 99.0, and that for the controls was 7.5. Sanford & Broadfoot(6) have drawn attention to the difficulties encountered in greenhouse tests for pathogenicity when done by the usual methods. Not much reliance was therefore placed on these tests, and the question of the relative pathogenicity of strains of *Fusarium culmorum* in England was left for more detailed investigation in the future.

Although isolation work was not commenced quite early enough, and was not done on a sufficient number of samples to give a complete picture of the activity of *F. culmorum* during the season, it is evident that the fungus must have been present in the soil from which these healthy wheat plants were taken, exerting no appreciable parasitic effect and entering the root systems, along with other fungi, only as the roots began to lose vitality after flowering of the crop. There was then a still further development of the fungus on the stubble, after the crop had been cut. This is in contrast to fairly numerous cases reported, especially from the north of England, of appreciable damage to wheat by the fungus *F. culmorum*. The soil or other factors which favour parasitic action by this fungus during the earlier stages in the growth of cereal plants are at present very imperfectly understood, but

it may be significant that these three instances in which no apparent injury was done were all on slightly alkaline clay soils.

The assistance of Miss E. Holmes and of Dr M. Fernando with portions of the cultural work, and of Dr W. A. R. Dillon Weston in the collection of samples from Cambridge, is gratefully acknowledged. Dr F. J. Greaney shared in the work described in the paper while he was a temporary worker at the Rothamsted Experimental Station. He is a member of the staff of the Dominion Rust Research Laboratory, Winnipeg, Canada.

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SYMPOSIUM AND DISCUSSION ON LABORATORY TECHNIQUE FOR EVALUATING FUNGICIDAL PROPERTIES

*Held on 18 January 1936, in the Botanical Department,
University College, Gower Street.*

The President, MR F. G. GOULD, in the Chair

I. PROTECTIVE FUNGICIDES AGAINST APPLE SCAB (*VENTURIA INAEQUALIS*)

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I. THE AIMS UNDERLYING THE WORK

VENTURIA INAEQUALIS was chosen as the test fungus because the need for the present investigation at East Malling arose out of field-spraying experiments designed to determine control measures for the disease caused by this fungus.

The test fungicide was Bordeaux mixture (proportion of copper sulphate to hydrated lime as 2 is to 3). This selection may be criticized in that this spray fluid is rather complex chemically and its action is not fully understood. Yet it was selected because of, rather than in spite of, that complexity. Something, at least, is known of its action and its properties, and if a laboratory method is evolved and built up by using a spray fluid of this nature, many of the difficulties inherent in the process will have been encountered and, in part, overcome.

The criterion selected was a simulation of field conditions as far as possible, not by exact reproduction of the many factors involved, but by standardization of such amongst them as lent themselves to standardization.

Hence, it was necessary to produce:

- (a) A standard surface on which to carry out the tests.
- (b) A standard, dry spray deposit, to which scab spores are added.
- (c) The thinnest complete film of spray fluid practicable (Serge Héranger⁽¹⁾ considers that $\frac{1}{10}$ mm. is the lower limit of thickness of film that can be obtained by spraying).
- (d) A standard process by which the spray deposit can be washed, as it is washed on leaves by rain, dew, etc.
- (e) A suspension of spores standardized for quantity and quality.

- (f) Standard conditions (temperature and moisture) for germination.
(g) A method of evaluating results.

II. METHODS OF PROCEDURE

(a) *Surface.* Apple leaves differ from one another even within a variety and they are not, at present, available all the year round. Glass slides, 3×1 in., chemically cleaned, are therefore used as offering the best available and convenient surface where action between the surface and the spray deposit will be infinitesimal or nil, and the influence of foreign matter can readily be eliminated. Three circles, each of exactly 15 mm. diam. (arbitrary), are scratched on each slide, by revolving the slide in a lathe in which a glass-cutter's diamond is mounted.

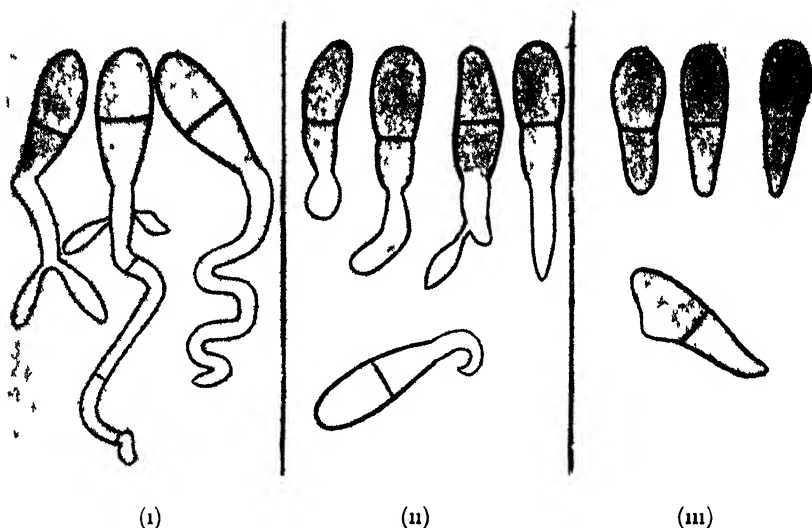
(b) and (c) *Spray fluid and the dry deposit.* Bordeaux mixture of known strength is applied to the area enclosed by each ring at the rate of 0.015 c.c. per ring. The fluid is delivered from a specially graduated, fine-bore glass pipette, and it is spread to the edge of the ring, where necessary, with a glass needle. The thickness of the fluid film is thus approximately $\frac{1}{10}$ mm. (*vide* Héranger⁽¹⁾). The slides are then air-dried and stored at 20° C. until next day.

(d) *Washing.* Rain water is variable in nature, and the use of running distilled water is extravagant and inconvenient to manipulate. The slides bearing the spray deposit are therefore placed face downwards on a glass rack in a bath of fresh distilled water, and rocked horizontally for given periods by mechanical means. Slides bearing a similar spray deposit are left unwashed as controls.

(e) and (f) *Spore suspension and incubation.* The fungus was cultured on ordinary malt-agar plates and on cheese-cloth wicks soaked in malt extract⁽²⁾, but neither method proved suitable for readily providing supplies of free spores (conidia). Eventually, cultures were developed on agar films and on pieces of sterilized apple wood, and both methods yielded free spores readily. Cultures are kept at constant temperature and with the minimum moisture necessary to prevent drying out. A spore suspension is usually obtained readily by washing the surface of the agar film, or of the piece of wood, with sterile distilled water. Nutriment tends to be extracted in varying amounts by the distilled water, both with agar films and apple-wood cultures. Methods of obviating this disadvantage have recently been tested. As a result, the use of agar films has been abandoned, for portions of the agar float in the spore suspension and cannot easily be removed. Suspensions obtained from apple-wood cultures can, however, be centrifuged and washed free from nutriment. After centrifuging, the supernatant liquid is poured off, sterile distilled water is

added to the concentrated spores, and the process is repeated until the spores ultimately remain suspended in pure distilled water.

It has been found necessary to obtain spore suspensions for tests from young cultures, for older cultures give mixtures of spores of different ages. It is now the practice to choose for the supply of spores cultures, 10–12 days old, which readily yield in suspension at least fifty spores per 2 mm. field. The suspension is ultimately regulated to contain about eighty spores per 2 mm. field. The spore suspension is put on the dry spray deposit on the slides at 0.04 cc. per ring by means of another specially graduated, fine-bore glass pipette, similar to that



Conidia of *Venturia inaequalis* showing the three groups into which the spores are classified when evaluating results of tests (i) germinated, (ii) inhibited after initial germination, (iii) ungerminated

used for the spray fluid. The same amount of spore suspension is also put within rings on slides without any spray deposit, and these act as controls and keep a check on the quality of the spores. The suspension is spread to the groove with a glass needle, and the slides are then placed in moist chambers (Petri dishes each containing a pool of distilled water to provide a saturated atmosphere) and incubated at constant temperature (20° C.).

(g) *Evaluation of results.* When the spores have germinated, but before growth has developed far enough to produce secondary conidia, germination is recorded by counting fields of spores more or less at random within each ring. The results are expressed at present in terms of spores (i) *germinated*—germ tube longer than spore, (ii) *inhibited after initial germination*—germ tube up to length of spore, (iii) *ungerminated*—no germ tube (see figure). It is usual on control slides to find at least 90 per cent of the spores in the first group.

III. RESULTS

Up to the present, the results obtained have been used to detect weaknesses in the method and thus to develop it, rather than to make comparisons of fungicidal efficiency. Many of the results could not reasonably be compared, for they were obtained under conditions that are now known to have been variable. However, certain points have emerged that are worth recording:

(a) Bordeaux mixture at very low concentrations is toxic to scab spores. A mixture containing as little as 0.0125 per cent copper sulphate (or, in practical terms, 2 oz. per 100 gallons of spray, which is one-sixty-fourth of the concentration commonly used in commercial spraying) virtually prevents germination even after the dry deposit has been washed continuously for at least an hour.

(b) The age of the culture, and the conditions of temperature and humidity under which it was grown, largely determine the quantity, and possibly the quality, of the spores readily liberated in distilled water. The later experiments have shown that there may also be certain other issues involved.

(c) The susceptibility of scab conidia to killing by Bordeaux mixture is governed by the presence or absence of nutriment in the suspension.

IV. FUTURE DEVELOPMENTS

It is anticipated that further work will be necessary before the method is perfected, while several of the tentative conclusions already reached need confirmation by more tests. The method will then be used for comparison of the fungicidal properties of new products (as yet not tested, or only partially tested, in the field) with those of the better-known sprays already in use commercially. Many points concerning the effect on fungicidal efficiency of the addition of spreaders, insecticides, or substances designed to obviate spray damage should, theoretically at least, be susceptible of evaluation by the use of this technique. It is often not easy to gauge by field trial the influence of these addenda on fungicidal value, for an apparently direct influence in these circumstances may be indirect, and due to differences in the physical properties of the spray fluids.

It is not unreasonable to hope that some light may be thrown on the fundamental action of certain spray fluids, either directly, or by leading up to a line of investigation through other channels. Certain questions with a possible bearing in that direction have already arisen during the present work with Bordeaux mixture.

Ultimately, promising new fungicides, which have proved non-phytotoxic in small trials in the field, will be tested on trees growing in pots under standard conditions in the greenhouse. This will involve the development of a satisfactory method of inoculating leaves

and fruits with the fungus. The final stage will be the intercomparison of the most promising new or improved spray fluids by field trial.

The continuation of this work, begun in November, 1934, was made possible by the aid of a grant from Imperial Chemical Industries, Ltd., to whom the writers record their indebtedness.

[Some of the apparatus used in this work was exhibited during the meeting, and the method of washing the spray deposit on the slides was demonstrated.]

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II. THE EVALUATION OF PROTECTIVE FUNGICIDES WITH SPECIAL REFERENCE TO APPLE SCAB *VENTURIA INAEQUALIS*¹

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III. PHYSICO-CHEMICAL ASPECTS

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THE status of physico-chemical methods in the laboratory examination of any particular fungicide is determined by the extent of knowledge of the mode of action of that and related substances. Once the fungicidal properties of the substance have been correlated with specific chemical and physical properties, laboratory testing is reduced to the evaluation of those properties. From the results obtained, the performance of the substance under field conditions can be foretold and an examination by biological methods is rendered unnecessary. For example, standardization by biological methods of lime sulphurs can be replaced by the analytical determination of content of polysulphide sulphur.

It is evident that the trustworthiness of the conclusions derived from the physico-chemical examination of the substance depends upon experience of the field performance of related compounds. For this reason, it is most important that the biological tests of any new product should have collateral physico-chemical tests designed not only for the purpose of defining the new product but for the eventual elucidation of its mode of action. The usefulness of such supplementary laboratory tests is illustrated by the following results obtained in an investigation of the protective fungicidal efficiency of

¹ See these *Transactions*, vol. xx, 304-9.

certain copper derivatives, Potato Blight being used as the test organism. As a working hypothesis it was assumed that the efficiency of the spray will be dependent upon the retention on the foliage of a copper-containing deposit from which the active fungicide is slowly formed. It follows that the protective efficiency is determined both by retention factors and by availability factors, and to separate these two sets of factors analytical studies were made of the sprayed potato foliage and the total amount of copper present at intervals after application was determined. Simultaneously, estimates of blight infestation were made in the field, with the results recorded in Table I.

The partial correlation between protective efficiency and retention is shown and differences attributable to availability can be determined. For example, comparing the relative efficiencies of cupric oxide and of cuprous oxide, the indications are that cupric oxide is better retained whereas cuprous oxide has the greater availability. In the dry

Table I

1934 trials			1935 trials		
Spray	Average estimate blight infestation	Average copper retention	Spray	Average estimate blight infestation	Average copper retention
Cotton-seed-Bordeaux	3.78	0.53	Cupric phosphate	4.40	0.38
Bordeaux	3.94	0.38	Cupric oxide	4.80	0.38
Cuprous cyanide	4.06	0.18	Cupric silicate	5.00	0.30
Cuprous oxide	4.44	0.18	Copper sulphite	5.20	0.12
Cupric oxide	5.33	0.22	Cupric oxychloride	5.53	0.20
Colloidal Burgundy	5.39	0.15	Burgundy Paste	5.43	0.19
Petroleum-Bordeaux	5.50	0.31	Cuprous oxide	5.50	0.19
			Cuprous cyanide	5.67	0.30
			Bordeaux	6.30	0.52
Significant difference	0.846	0.05		0.318	—

season of 1934 the latter factor was predominant and cuprous oxide proved the superior protector; in the wet 1935 season, the greater amount of cupric oxide retained overshadowed its lower availability and cupric oxide proved the better fungicide. Conclusions of this character must obviously be confirmed by further trial and they are instanced only for the purpose of demonstrating the value of physico-chemical tests as supplements to field trials. It is not, at this stage, suggested that they are essential and it may be possible to subject an unknown product *XYZ* to field trials and, if found satisfactory, to admit the product to the category of fungicides suitable for recommendation. Physico-chemical criteria play no part in such a scheme but, once the field trials are replaced by laboratory biological tests, physico-chemical considerations again require attention.

The simplest definition of a laboratory trial is one in which one or more of the factors affecting fungicidal efficiency which are variable in the field trial are maintained constant. This definition may be

illustrated by a brief description of Salmon's method of assessing direct fungicidal properties, which employs the hop Powdery Mildew as the test fungus. By using hop plants vegetatively propagated from one parent plant susceptible to the mildew and by selecting for treatment leaves and mildew patches in the same stages of active growth, the biological condition of the fungus and host plant is standardized and variations in their condition which may affect the response of the fungus to the fungicide are diminished. Further, the physical properties of the spray are modified so that it is able to displace air from and to wet the conidiophore mass. Finally, a sufficient amount of spray is applied to achieve complete wetting. In interpreting to practice the results obtained by this technique, it must be remembered that, as variations in the biological condition of the fungus and host plant have been eliminated, the results are of relative and not absolute value. Further, that the results are applicable only to sprays which have wetting properties sufficient to wet the conidiophores when applied in heavy amounts in the field.

The accuracy of the interpretation of the results of laboratory tests is dependent upon the correctness of the allowances made for the influence of variations in the factors held constant in the trial. This theme has been developed in a previous discussion⁽¹⁾ and it is only necessary to emphasize that the missing variables are always found to involve physico-chemical factors. The neglect of such criteria, which was conceivable in field trials, becomes highly dangerous in laboratory tests.

From the biological aspect, the laboratory trial is generally designed for the examination of the toxicity of the material to fungi. Fungicidal efficiency, although primarily dependent upon actual toxicity, is determined by other factors, of which the amount of material applied and retained by the fungus or plant surface may be selected as an example. In all laboratory methods of evaluating fungicidal properties, standard procedure is adopted for the application of the spray or dust under test. Moore and Montgomery spread a definite volume of spray over a standard area of surface; Marsh and Salmon both apply the spray under standard conditions to the surface held at right angles to the direction of the spray; Marsh spraying for a definite time, Salmon spraying copious amounts of a spray of good wetting properties. What assumptions do these various procedures involve and how are the results comparable with those of practical spray methods?

With the surface held at right angles to the direction of the spray it has been found⁽²⁾ that, with solid-liquid systems akin to those encountered in spray practice, the whole of the spray applied is retained up to a certain amount dependent upon the particular liquid-solid system. Irrespective of the spray or surface, the amount re-

tained is equal to the amount applied until the volume present is such that the excess runs or drains off the surface. By the technique used by Marsh, therefore, standard amounts of spray are applied per unit area provided that the time of spraying be so selected that no run-off or drain-off of the spray occurs. The technique thus produces the same end-point as that of Moore and Montgomery except that the latter method makes no apparent provision for the possibility that excess of spray may be applied to the surface. In Salmon's method, excess of spray is deliberately applied and the amount retained will be determined by the wetting properties of the particular spray under test. But a requisite of the technique is that the wetting properties of the spray shall be sufficient to permit complete wetting. Under such conditions it has been shown that approximately constant amounts of spray are retained.

Salmon was concerned with direct fungicidal properties for which purpose only sprays of good wetting properties are suitable. The other techniques under discussion are applied to sprays of indifferent wetting properties such as are normally used as protective fungicides. With such sprays, the amounts retained upon foliage will be limited by the maximum amount retainable, an amount which has been shown to be greatly influenced by the physical properties of the spray. Thus upon a cellulose nitrate surface, the maximum retainable weight of water is of the order of 200 mg. per sq. in.; that of a solution of an effective wetting agent is of the order of 25 mg. per sq. in.

There is, however, a compensating set of factors, for the condition that the surface shall be held perpendicular to the direction of the spray is but rarely met in practice. If the surface be tilted to the spray it is found that only a proportion is retained, the proportion increasing with the wetting properties of the spray. When applied in limited amount, therefore, a spray of wetting properties similar to those of water will be retained in less amount than a spray of good wetting properties provided that, in neither case, does drip occur.

These considerations concern simple aqueous solutions, and with heterogeneous sprays, such as suspensions and emulsions, other physical factors come into play. Enough has been said to show the complexity of the apparently simple problem of how to apply the spray in the laboratory test. The exact reproduction of the spray deposit obtained in practice is not necessary provided that the effect of the factors held constant in the laboratory test be remembered in translating the results to practice. For this reason there is much to be said for the simplest technique which will give consistent results and of which the significance in relation to field technique is understood.

The method of application is but one example of the physico-chemical considerations involved in laboratory tests, but it is enough to illustrate the contention that, if the results of biological toxicity

trials are to be applied to practice, consideration be paid to the physico-chemical properties of the spray or dust.

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IV. FUNGICIDES SUITABLE FOR FOOD PRESERVATION

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INNUMERABLE substances possess fungicidal properties. Whether or not any of these substances can be usefully employed to prevent the growth of fungi in any particular instance depends not on their toxicity alone, but also on other properties such as solubility, taste, colour and smell. Consider, for example, the fungicides which are employed in a variety of ways in food preservation. Hypochlorites and formalin are used to sterilize storage rooms and grading machines and packing sheds. Benzoic acid or sulphur dioxide is added to certain foods to preserve them. Oranges are sometimes dipped in a solution of borax or sodium bicarbonate in order to reduce the incidence of rotting by green rot. Carbon dioxide is sometimes added to ships' holds to delay the onset of mouldiness and sliminess of meat carried in them. It has been suggested that certain fungicides might with advantage be added to the paper in which fruit is wrapped to prevent the onset or the spread of rotting. The compounds used in these various ways all possess the property of retarding or preventing growth, but otherwise the properties which make them useful are very different.

Because usefulness depends on the combined effects of a variety of factors, it is generally agreed that the advantage of using a fungicide in any particular instance can be tested only by direct experiment. Preliminary laboratory tests of substances can be undertaken to find:

- (1) Their action on germination and growth of fungi or their power of killing, i.e. their fungicidal action.
- (2) Their solubility, volatility, and other properties which may be of importance.

From the results of such tests, it is usually possible to decide what substances are obviously unsuitable and what substances may be suitable for further trial for the purpose for which they are wanted.

LABORATORY TECHNIQUE FOR EVALUATING FUNGICIDAL PROPERTIES

The type of test undertaken to find how fungicidal a substance is, is usually of the simplest order. This is to be regretted, for certain qualitative differences in the manner in which growth is re-

duced in the presence of inhibitors are often of importance in determining choice of fungicides.

To discover the effect of a substance on growth, it is necessary to measure germination or growth in the presence of increasing amounts of the substance. The difficulties in doing this are twofold.

There is the difficulty of knowing what criteria of germination or growth to adopt; and there is the difficulty of technique of carrying out the experiment.

The difficulty of knowing what criteria of germination or growth to adopt is due to the fact that substances may affect the course of germination by:

- (i) decreasing the number of spores which germinate,
 - (ii) increasing or decreasing the latent period,
 - (iii) decreasing the rate of elongation of the germ tubes;
- and the course of growth by:

- (i) decreasing the rate of radial spread,
- (ii) decreasing the density of the mycelium,
- (iii) altering the final yield.

Moreover, these aspects of germination and growth may be affected differently as can be seen when one compares the effect, for example, of temperature, alcohol, carbon dioxide, oxygen and carbon monoxide on growth. For these reasons, any single criterion of growth is not sufficient to give adequate information about the action of a fungicide.

No one, I think, would suggest that the effect of temperature on the germination and growth of a fungus was fully described if the temperature at which 50 per cent of a sample of spores of the fungus failed to germinate was given. Yet one frequently finds some such value as the concentration of a substance needed to inhibit the germination of 50 per cent of a sample of spores claimed as a measure sufficient to describe the action of a fungicide. These points have been considered before⁽¹⁾.

The difficulties of technique include the choice of medium and the maintenance of the required condition. The following method has been found most suitable for studying the effects of soluble non-volatile inhibitors.

A sheet of squared paper is placed on a smooth level bench in a constant-temperature room, and covered with a sheet of glass. Petri dishes, 9 cm. in diameter, are set out in regular order on the glass, and into them are placed 10, 9.9, 9.8, 9.7, etc. c.c. of a liquid nutrient medium (2 per cent malt). To the dishes are then added 0, 0.1, 0.2, 0.3, etc. c.c. of the solution of the fungicide, ten to twenty times as strong as that necessary to prevent growth. (Preliminary experiments may be necessary to find out what concentration of fungicide is required to prevent growth.) In this way, a series of solutions containing increasing amounts of the substance is obtained.

Into the centre of each dish is placed a small piece of porous pot, previously sterilized and then inoculated with a small drop of a suspension of spores of the fungus. The spores germinate and the hyphae grow out regularly in every direction to form a circular mycelial mat, just as when growth is on a solid agar medium. The increase in size of the diameter of the colony can be measured by the aid of the squared paper. Care is required, especially in the early stage of growth, in examining and measuring the colonies, or the tips of the hyphae are thrown out of position and growth becomes irregular.

It is not always possible to make solutions ten to twenty times as strong as those which inhibit growth, and in such instances, some slight modifications are needed in making up the culture solutions in the dishes. For example 5 c.c. of a stronger culture medium may be added to each dish, and then 0, 1, 2, 3, 4, 5 c.c. of fungicidal solution and 5, 4, 3, 2, 1, 0 c.c. of water to make up to 10 c.c. in each dish.

The advantages of using liquid cultures are many. The solutions can be made up from two stock solutions sterilized separately. If there is any danger of the solutions in any of the Petri dishes changing in composition or *pH*, owing to the growth of the fungus, they can be sucked off daily and replaced by new solutions of the required composition. What is more, the composition of the solution can be changed after growth has started. It is usually advisable to buffer the nutrient solution with phosphate or phthalate buffers. When the diameter is plotted against time, a straight line is obtained. The slope of this line—the rate of increase in size of diameter—is generally the most suitable measure of the rate of growth.

HOW THE MANNER IN WHICH A FUNGICIDE RETARDS GROWTHS MAY AFFECT ITS VALUE AS A MEANS OF PRESERVING FOOD MATERIALS

Graph I shows the effect of temperature and Graph II the effect of alcohol on the growth of a fungus.

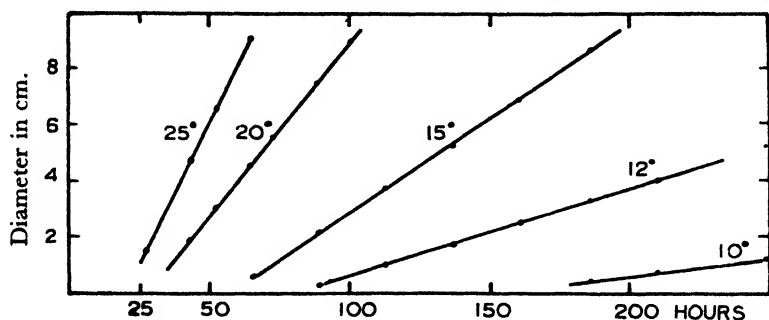
Lowering the temperature has two effects. The germination and early growth is delayed, and the rate of subsequent growth is decreased. The presence of alcohol decreases the rate of growth, but does so without any, or only a slight delay of germination.

The action of some substances (CO_2 and NH_3) is somewhat similar to that of temperature. The action of other substances (SO_2 and benzoic acid) is similar to that of alcohol.

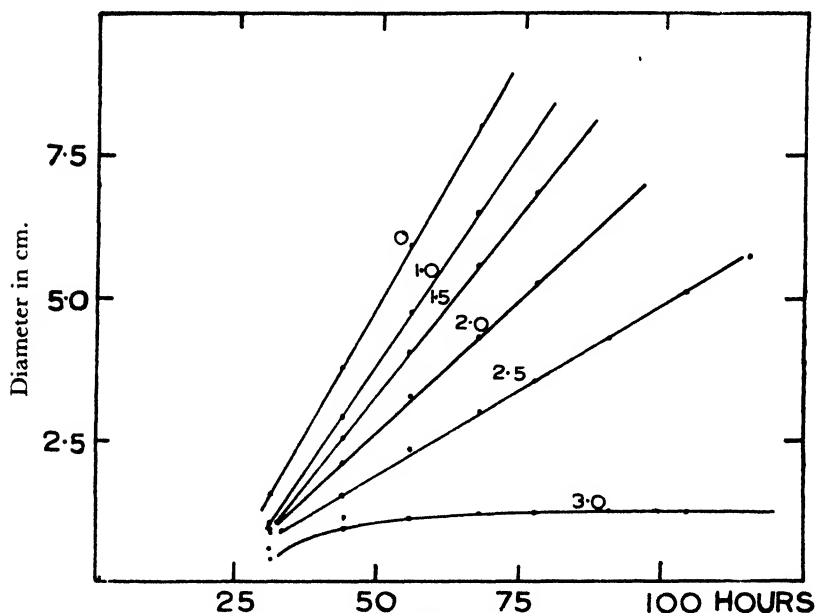
Thus the time taken for a fungus to grow to a definite size is altered to various extents by the presence of different inhibitors. The importance of this in food preservation is as follows: When food is stored, the length of time it can be kept (storage life) is of first importance. The time for food to become obviously mouldy is in certain

instances about the same as that required for the fungus to grow to 1 cm. in diameter, on a nutrient agar medium kept under the same conditions as the food. Now, whereas by decreasing the temperature from 25 to 10° the time required for the colony to grow to 1 cm. in diameter is increased from 25 to 240 hours, the effect of adding alcohol is to increase the time from 25 to not more than 40 hours.

By lowering the temperature sufficiently, or by raising the concentration of alcohol sufficiently, growth can be inhibited completely,



Graph I. Growth of *Trichoderma* at different temperatures

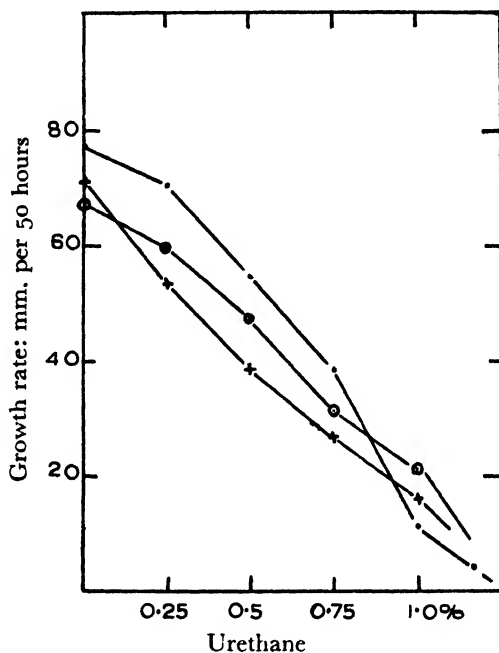


Graph II. Growth of *Trichoderma* in the presence of alcohol. (Parts per 100)

and the effect as far as mould growth is concerned is the same. But, except when conditions are so arranged completely to retard growth, the advantage of using low temperature or substances having a similar action, rather than alcohol (or one having a similar action) is obvious—though generally unrecognized.

THE EFFECT OF THE MEDIUM ON THE FUNGICIDAL
PROPERTIES OF SUBSTANCES

The fungicidal properties of a substance often depend on temperature and the medium in which it is acting. One factor which often has a marked effect is the acidity of the medium.



Graph III. Growth of *Trichoderma*. \odot pH 4.0, \cdot pH 5.0, $+$ pH 6.0

The influence of pH on the action of three inhibitors is illustrated in Graphs III, IV and V, in which the rate of growth (as measured by rate of linear spread) is plotted against the concentration of the inhibitor.

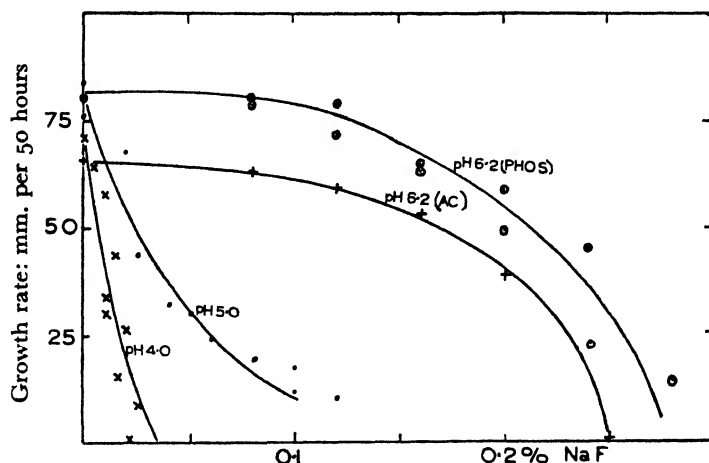
Graph III shows that the retarding effect of urethane on growth is directly proportional to the concentration and within wide limits unaffected by the pH of the medium.

Graph IV shows that the retarding effect of sodium fluoride is much more pronounced in acid than in alkaline solution, and that the relation between concentration and inhibition also varies with the alteration in the acidity of the medium.

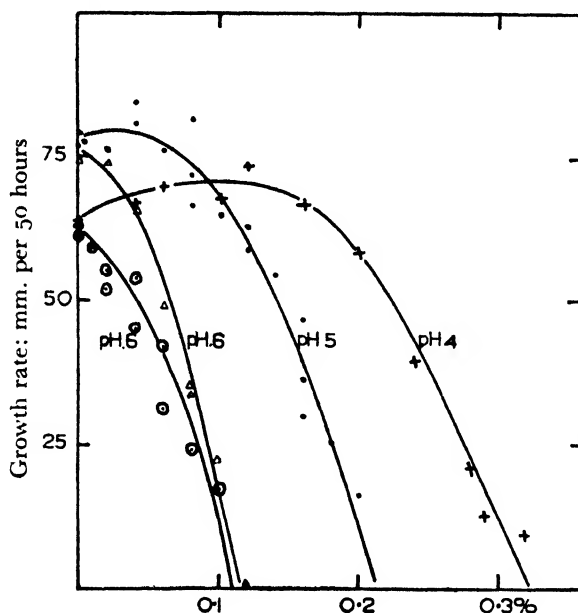
Graph V shows that the retarding effect of borax is more pronounced in alkaline than in acid solutions, and that the relation between concentration and inhibition varies as the acidity of the medium alters.

These figures suggest that in choosing a fungicide, some consideration of the acidity of the medium in which it is to be used is necessary,

and also that by suitably modifying the acidity of the medium, the efficiency of some fungicides can be increased. For example, when a substance such as urethane is used, no consideration need be paid to



Graph IV. Growth of *Trichoderma* in presence of fluoride



Graph V. Growth of *Trichoderma* in the presence of borax

acidity. But it might be profitable when sodium fluoride is used to make the medium more acid and use a smaller concentration of fluoride, than use a higher concentration of fluoride while leaving the acidity unchanged. Similarly when borax is used it might be profitable to make the solution more alkaline and use a lower concentra-

tion of borax, rather than increase the strength of borax and leave the acidity of the medium unchanged.

These simple examples show that detailed knowledge of fungicidal properties often suggests ways of improving fungicidal action.

However, when one tries to discover from published results how the action of fungicides is affected by various factors, one finds, strange though it may sound, that the details of the action of very few substances are at present known.

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V THE LABORATORY TESTING OF WOOD PRESERVATIVES

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THE laboratory tests upon wood preservatives have been mainly of three types:

- (1) Tests of toxicity in artificial agar media.
- (2) Tests of toxicity in wood blocks.
- (3) Tests of permanence, e.g. volatility and leaching tests on treated wood blocks.

In this discussion only the first two types of tests are referred to.

In the past so-called agar or Petri-dish tests have been extensively used, particularly in the U.S.A., for evaluating the toxicity of wood preservatives. They give clear-cut results which can be repeated if carried out under standard conditions. Continental workers have criticized the use of agar medium and have developed the wood-block test. Results obtained with experiments on wood are liable to vary and the experimental error may be high, but results in agar tests which may appear to be concordant can lead to very erroneous conclusions, since the reactions between an antiseptic with agar and with wood may be utterly different.

We now regard the agar test solely as a preliminary sorting-out test. It is useful for comparing the toxicities of substances of generally similar chemical composition, e.g. of various phenols, but is quite unsuitable for comparing the toxicity of substances of widely different composition, e.g. of creosote and of sodium arsenate. In carrying out tests in agar it is most important that the preservative and the medium should not be sterilized after mixing. Sterilized amounts of medium should be added aseptically to measured amounts of the preservative.

The total amount of nutritive medium and the total volume must be kept constant at each concentration. The end-point or toxic point should be expressed as the interval between that concentration at which growth just takes place and the concentration next above it in the series at which no growth takes place.

Except for such sorting-out tests, it is essential to use wood as a substratum for tests with wood preservatives. The species of wood used should be one which is readily decayed and has a low natural resistance to the attack of wood-destroying fungi. It should also be one which can easily be impregnated with a preservative. The sapwood of Scots Pine, *Pinus sylvestris*, and the outer wood of beech, *Fagus sylvatica*, are suitable woods.

The most useful figure for comparing the efficiency of wood preservatives is the minimum quantity of preservative which prevents all attack by certain organisms. This toxic point should be expressed as amount of preservative per unit volume of wood; it is useful also to give the strength of solution used to attain this concentration in the wood. The method of applying the preservative must be such that there is a uniform distribution of antiseptic. The most satisfactory method of achieving this is to impregnate thoroughly under vacuum, the blocks which previously have been sterilized and dried in an oven at 100° C., the samples being weighed before and after treatment to determine the absorptions obtained. Water soluble substances are applied in aqueous solution, oily materials in a suitable solvent such as petroleum-ether or acetone. After treatment with an aqueous solution the moisture content of the samples must be reduced to the point most suitable for fungal attack, i.e. to about 40 per cent of the oven-dry weight. Where an organic solvent has been used this must be completely evaporated away. This is accomplished by drying the blocks in an oven provided with forced ventilation at a temperature not exceeding 50° C.

The blocks are exposed to infection by placing them in contact with cultures of certain wood-destroying fungi growing on agar medium in special culture flasks. They are incubated for four months at 22° C. and kept in a moist condition by the addition of water to a reservoir in the neck of the culture flasks. At the end of this period the blocks are removed from the cultures, and examined for signs of fungal growth. They are freed from any superficial mycelium, then oven-dried and re-weighed. An indication of the amount of decay if any, which has occurred, is given in the loss in dry weight (initial dry weight – final dry weight), which is then expressed as a percentage of the original dry weight; allowance being made where necessary for the weight of preservative absorbed. In carrying out wood-block tests reliable results may be obtained but precise results cannot be expected.

A number of test fungi must be used since the wood-destroying fungi are not uniformly resistant to antiseptics. A fungus may be resistant to one substance and sensitive to another, for instance *Fomes annosus* is resistant to creosote and sensitive to zinc chloride, *Coniophora cerebella* is resistant to zinc chloride and very sensitive to creosote. At least three test fungi must be used, these should be:

- (1) Active wood destroyers.
- (2) Resistant to fungicides.
- (3) Not unduly sensitive to slight variations in environmental conditions.

Lentinus lepideus Fr., *Coniophora cerebella* Pers. and *Poria vaporaria* (Pers.) Fr., are suitable for tests on pine, and *Polystictus versicolor* (Linn.) Fr. for tests on beech.

Summarizing the important points to be considered in making laboratory tests upon the toxicity of antiseptics for the preservation of materials:

(I) The tests should be made with pure cultures of the organisms against which it is desired to protect the material. With wood preservatives a number of test fungi should be used.

(II) The medium used must be the material which the anti-septic is designed to protect, in as nearly as possible its normal physical condition.

(III) Conditions for the growth of the injurious organism should be optimum.

VI. SEED PROTECTION

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IN horticultural and agricultural practice the fungicides in use may be divided into three major groups: (a) fungicidal solutions, emulsions or suspensions, (b) fungicidal gases, (c) fungicidal dusts. It would be difficult, and probably impossible, to devise any one method of testing for such diverse phases. Although it is relatively easy to test simple soluble inorganic and organic solutions against a given fungal spore or spores when these are not in contact with their hosts, it cannot be inferred that the values so obtained will produce corresponding values in the field.

It would seem clear that laboratory technique for the evaluation of fungicidal values should be regarded as a rather close-meshed sieve to sort out probable fungicides from improbable ones. In such a sifting it may well be that evidence will be obtained that will indicate

relationships between chemical and physical properties, and fungicidal values and phytocidal effects.

It is the purpose here to discuss particular types of fungicides, viz. those that are used for the disinfection of seed. The disinfection of grain to prevent seed-borne disease is a well-established practice of plant hygiene and needs no comment. The methods employed, however, may alter. In recent years many farmers have changed from the wet treatments to dry, the latter consisting of disinfectant dusts that contain salts of mercury as their fungicidal basis. These dusts—complicated in chemical and physical properties—are now widely used, and by many are considered to be an advance on the older wet treatment, since, when correctly applied, they give control of Leaf Spot of oats and Leaf Stripe of barley—two diseases that were not adequately controlled by a wet treatment such as a solution of the gas formaldehyde in water. Moreover they are of value in the treatment of certain seeds other than cereals and better stands are sometimes established by the use of such treated seed. It is here that a slight digression is necessary. Laboratory tests are specific against a given organism but the seed when planted is in an environment in which it is in competition with a large and variable soil flora and fauna. To be effective the fungicide used must be capable of killing any disease organism adhering to or slightly penetrating into the seed, and also confer upon it *subsequent protection*. It is the analogy of Listerian practice that was founded on Pasteurian principle.

At Cambridge an investigation has been in progress for some years on seed protection, particularly on disinfectant dusts that contain salts of mercury, and a large range of these has been tested. As such dusts contain not only a fungicidal salt but a carrier for this, it is clear that *at least* two phases must be considered, i.e. the salt and the filler, and the chemical and physical properties of both must be known. Since these are two variables it does not follow that if a fungicide *A* has a greater value than a fungicide *B*, that a dust containing *A* is more potent than *B*.

Although theoretically—knowing the high toxicity of the salt—one might assume that a dust containing 5 per cent of mercuric chloride would be a good seed disinfectant (if it were practical to use it), this is not necessarily so, for when such a dust was used to treat infected oats the reduction in the infection of Leaf Spot was from 24 to 12 per cent only. The control, however, with Bunt contaminated wheat was better, the reduction being from 21 to 5 per cent.

Somewhat similar parallels could be drawn from a large range of inorganic mercury salts that we have tested. In this connexion too it may be of interest to record that when spores of *Tilletia* were exposed to the vapour of ethyl mercuric iodide this appeared to exert a lethal action. When, however, a dust containing ethyl mercuric iodide was

used on Bunt contaminated wheat sown in the field there was very little control of the disease, although the same dust gave a good control of Leaf Spot of oats. The sodium salt of hydroxy-mercuridibrom-fluorescein, although a well-known bactericide, gave little or no control of Leaf Spot of oats.

Observations such as these suggest that although some of the inorganic and organic mercury salts are specific for certain seed-borne diseases, it does not follow that they are a panacea for all of them; moreover they indicate that laboratory evaluations, if unsupported by small-scale field tests, may be misleading.

It was for such reasons as these that we have held the intrinsic test for fungicidal values to be the disease control that is obtained on small-scale replicated and randomized field plots. Although biological laboratory work has served our purpose it has been in the field that our most useful records have been obtained. In this type of work a large laboratory garden is essential, and in the early part of our investigation we relied exclusively on this. In our preliminary trials we used the following technique. Dusts were prepared containing most of the known inorganic compounds of mercury and a selection from the organic compounds and these were compounded with the same filler. 5 and 10 per cent of the appropriate salts was used in these dusts and they were employed to control Leaf Spot of oats. The dose was at the rate of 2 oz. to the bushel of grain and the dusting operation was carried out for a period of 1 min. Each plot consisted of a rod row and each of these was replicated five times. Certain proprietary dusts were included for the purpose of comparison.

As the resulting estimates were made by recording the disease present in two hundred oat seedlings per plot there was therefore a total count of one thousand per treatment. In these preliminary experiments the plots were not randomized, but each treatment was spaced at regular intervals throughout the field and every tenth plot was an untreated one. Although this procedure has varied slightly a somewhat similar one has been adopted when investigating the control of other diseases and also in the various problems that have arisen during the course of the work. In several of the later trials eight replicates have been made and these have been randomized. In the main trials we have tried to assess the relative fungicidal values on a total count of not less than one thousand seedlings or ears per treatment. The results from some of these experiments we have already given⁽¹⁾.

The laboratory investigation was concerned with solutions of pure salts at known concentrations—in particular those that could be represented by the general formula $R\text{-Hg-X}$ where R was a hydrocarbon and X an acidic radical. While at first sight the number of possible variants of R appears great, practical limitations such as

stability and methods of synthesis reduce the number to the following four:

Tolyl	C_7H_7	Ethyl	C_2H_5
Phenyl	C_6H_5	Methyl	CH_3 .

This work was not extensive and we made no attempt to devise new methods to test existing seed disinfectants or to elaborate technique for a laboratory testing of fungicidal values. The objects of this investigation were primarily to obtain evidence on the relationship of toxic action to chemical constitution and to obtain information on the phytocidal effect of certain salts. The method we used was a simple one. In effect it was to determine the minimum concentration of the compound that would give the maximum control of the disease organism, i.e. *dosis curativa*, and to determine then the maximum concentration that the seed could tolerate, i.e. *dosis tolerata*. The latter was obtained by germination tests carried out at the Official Seed Testing Station, Cambridge. In our preliminary experiments watch-glasses containing Dox's medium were lightly flooded with a suspension of the spores of *Tilletia caries* and were exposed for varying periods to the vapours of the particular compounds. The amount of the spore suspension used and the period of exposure to the vapours was constant for a particular series of experiments. Rapid drying out of the medium was retarded by superimposing watch-glasses lined with unsized and moistened paper. Later, the percentage kill was determined by microscopic examination; and, subsequently, natural fungal and bacterial contaminations that developed on the treated media were noted, and the extent of these assessed and recorded.

Somewhat similar tests were made with solutions, chiefly with those containing methyl compounds. Spores of different species of *Ustilago* were used, since these germinate within twenty-four hours, whereas those of *Tilletia* take several days. A suspension of these in distilled water was added to glasses containing either Dox's or potato agar medium and the appropriate salt solution at a known concentration was then added.

A range of salts was also tested against the organism causing black-arm of cotton, *Bacterium malvacearum*. The technique employed was to take 5 c.c. of bouillon and glycerine, add 5 c.c. of a suspension of the organism in distilled water and then 5 c.c. of the appropriate concentration of the fungicide concerned. The point at which the organism was inhibited or killed was determined by the turbidity of the suspension or by reinoculation into sterile bouillon. The bactericidal activity of some of the compounds was also made by Rideal-Walker carbolic coefficient tests.

As some of the compounds we are using are volatile and are fungicidal in very small quantities, we are now trying to devise some method for estimating the relative fungicidal value of minute amounts

of them. The method that we are at present experimenting with is as follows: A small bulk of seed is dusted with a given quantity of the compound. The number of seeds per pound is known, consequently the amount of chemical per seed can be estimated. In this way minute quantities of the compound can be obtained and tested. A number of seeds so treated are then placed in a litre bottle in which is a small beaker containing a given quantity of water. A glass slide treated with potato agar on which spores of *Ustilago* are present is then suspended from the cork that seals the bottle. After twenty-four hours the glass slides are examined to see if the spores have been killed.

I have made no reference here to the results that have been obtained by these methods of investigation, or to those qualities—other than fungicidal efficiency—that are desirable in modern seed disinfectants, as it may be that such factors do not come within the scope of this symposium.

REFERENCE

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VII. SEED PROTECTION

By A. E. MUSKETT

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FROM 1929 to 1931 a number of experiments was carried out in Northern Ireland with the object of determining the most satisfactory treatment for securing the control of Smut diseases in the oat crop. Among the materials used for the purpose of seed disinfection were proprietary organic mercurials which at that time had not been placed on the market. The results obtained from these experiments showed that whereas formaldehyde gave almost perfect control of Smut diseases, the organic mercury compounds used were superior as general disinfectants against seed-borne diseases. As, at the same time, these compounds proved to be very little inferior to formaldehyde in securing Smut control and as there appeared to be no risk of seed injury attendant upon their use, it seemed probable that there was every likelihood of their being of considerable value in agricultural practice. To-day the use of these compounds is becoming widespread with the result that the plant pathologist whose duty it is to advise as to the most suitable measures to employ in securing the prevention of crop diseases is faced with one more problem, viz. he must be able to distinguish clearly between the products which are efficient and those

which are not. For the time being at any rate the advent of this new type of proprietary seed disinfectant has complicated the issue, and to put the matter bluntly, an improvement in the methods (or the devising of new methods) by means of which the values of fungicidal materials may be assessed as rapidly and accurately as possible is urgently needed. Biological technique only has been the subject of investigations in Northern Ireland and no attempt has been made to approach the question from chemical and physical standpoints. The object of the work has been to assess accurately the value of a fungicide by testing it against the disease which it is desired to prevent.

Helminthosporium disease of oats is common in Northern Ireland, and it was decided to attempt to devise a suitable technique for the testing of fungicides for the prevention of this disease. The problem was first approached by carrying out carefully planned field trials. Towards this end, seed known to be heavily infected with *Helminthosporium* was sown in drills at a standard rate at the normal sowing time, and when the seedlings had reached the optimum stage of development they were examined for the incidence of the primary phase of the disease. The results obtained were useful but lacked precision, and it became clear that several seasons' work would be necessary before really accurate data could be obtained. Furthermore, other factors arose which indicated that a more precise method applied under carefully controlled conditions was highly desirable in addition to field trials. In brief these factors may be cited as follows:

(a) A large number of treatments were used and it was found impracticable to lay down the number of plots necessary to ensure the statistical accuracy of the results obtained. The employment of an accurate small-scale technique would at least allow a number of inefficient materials to be discarded in the preliminary stages.

(b) It is important that the counting of diseased seedlings be concluded before the onset of the secondary phase of the disease. The undertaking of field experiments on too large a scale makes this difficult, while a period of inclement weather during the critical period may impose a further difficulty.

(c) The extent of the incidence of the disease varies largely from season to season in accordance with the weather conditions experienced.

Before attempting to devise a small-scale method of testing it was decided to adopt some definition of the ideal seed disinfectant and the following was chosen:

The ideal seed disinfectant shall be that which will secure complete control of a seed-borne disease *when the crop is grown from heavily infected seed under conditions designed to produce the maximum incidence of disease*. It will exert no adverse effect upon the crop, be non-poisonous

to animals or man, and its use will entail a minimum expenditure of money and labour.

The adoption of the conditions laid down by this definition involved a considerable amount of preliminary work in order to determine the conditions under which seed oats heavily infected with *Helminthosporium* will produce a crop showing the maximum incidence of disease. The seedlings were raised in round cake tins filled with soil and each capable of accommodating a hundred plants. An equal quantity of soil was weighed into each tin and the seeds were sown at a constant depth and equally spaced. Each seed was dropped from forceps into the hole prepared to receive it in order that the sowing process should remove as little of the disinfectant as possible. After sowing, the same weight of covering soil was used for each tin. The plants were grown to such a stage as to ensure that the primary phase of the disease had developed to its fullest extent in the crop. By means of this method the influence of soil, moisture, and temperature upon the incidence of the primary phase of the disease in the crop was investigated. Two types of soil were used, one light and sandy, the other a heavy loam. As no large greenhouse with adequate temperature control was available, the effect of temperature was studied by making five different sowings during the period from 1 March to 1 May, which represents the extreme sowing range for the spring oat crop. Two recording thermographs were run in conjunction with the experiment and an average temperature for each sowing was calculated from records made at intervals of two hours throughout the experimental period. Three degrees of soil moisture were used, one representing a very wet soil, the second a soil with a medium moisture content and the third a very dry soil. The access of rain to the tins was prevented by carrying out the work in an outside verandah built against a wall and with a roof sufficiently wide to prevent the rain from entering. In order to ensure the constant moisture content of the soil in the tins during the experimental period, each tin was watered, every other day, until its weight was the same as that recorded at the commencement of the experiment. The time of the experiment varied for the various sowings from sixty to thirty days according to the temperature and moisture conditions prevailing.

In brief, the results from these experiments have shown that those conditions which favour the good growth of the oat crop lead to a minimum incidence of *Helminthosporium* disease. The pre-emergence phase of the disease is pronounced under conditions which make for the slow germination of the seed and growth of the seedling; it was particularly marked when low temperature conditions prevailed. The minimum incidence of disease occurred under conditions of wet soil and high temperature. In dry soil the disease was severe even under

conditions of high temperature. Soil type is probably the least important of the three factors investigated, light soils tending to favour the incidence of the disease more than heavy soils.

For testing purposes a good medium loam is satisfactory. The temperature and moisture conditions should be such as will lead to a maximum incidence of the post-emergence phase of the disease and a minimum of the pre-emergence phase. An average temperature of from 8 to 9° C. throughout the experiment provides those conditions, and in Northern Ireland for the two seasons 1934 and 1935 they have been obtained by commencing the test during the second week of March. As wet soil does not encourage the development of the disease to its fullest extent, such soil is not used for testing purposes. Both moderately moist soil and dry soil are used, the moderately moist because it approaches natural conditions and the dry because it has been found that dust disinfectants tend to be less effective under dry soil conditions, which must therefore be included so that the efficiency of any disinfectant can be correctly assessed.

These necessary conditions for testing were first investigated in the spring of 1934 and results obtained from repeat experiments made in 1935 fully confirmed the findings. A series of comparative tests using a large number of fungicides was carried out in 1935 and very satisfactory results were obtained.

CONTINENTAL METHODS

In the summer of 1934 I visited the Continent and was privileged to inspect the work being carried out at the State Biological Institute, Berlin, in connexion with the official testing of fungicides and insecticides. The following is a summary of some of the methods and technique employed with regard to the testing of seed disinfectants.

The German Plant Protection Service publishes an official list of approved plant protection substances. To qualify for inclusion in this list a material must satisfactorily pass a series of official tests. Materials which can be prepared by any person following given instructions or to the use of which objection may be taken in the public interest, cannot be included. The tests to which materials are subjected fall into two groups: (i) The Preliminary Tests, and (ii) The Main Tests.

THE PRELIMINARY TESTS

These are carried out at three of the central stations in Germany, the manufacturer being allowed the choice of station.

The *seed grain used for the tests* must have a normal water content (13–14 per cent). For Bunt, clean seed wheat is used and is artificially infected with *Tilletia* spores at the rate of 0.2 g. per 100 g. of grain.

For Loose Smut in oats two seed samples from different sources are used. At least one of these samples is taken from a crop the plants of which were infected at flowering by dusting them with spores of *Ustilago Avenae*. Seed rye must not be infected with common moulds to an extent of more than 5 per cent.

Germination tests are carried out according to the methods employed at the official seed-testing station. The samples used for this purpose must be the same as those intended for use in greenhouse and field trials. Special attention is paid to the question of seed injury and where this is suspected soil tests are carried out in addition. Tests carried out at temperatures of from 6 to 10° C. are made when possible, in addition to those made at from 18 to 22° C.

Laboratory methods used for the determination of fungicidal efficiency in the case of Bunt in wheat vary according to the method of disinfection employed. For the *steeping* process *Tilletia* spores are directly treated with the fungicide and then tested for viability at stated intervals after they have been plated out on moist soil contained in Petri dishes. Tests for the *sprinkling*, *short-steeping* and *dusting* processes are made by treating artificially infected grain which is then sown on the surface of moistened soil similar to that used for the steeping process. The *Tilletia* spores present are examined for viability at stated intervals. The methods used for Stripe diseases of cereals caused by *Helminthosporium* spp. are similar to those employed for Bunt in wheat. For *Fusarium* disease of rye, tests are made in the *greenhouse* by planting disinfected seed in soil in wooden boxes. The spread of the mycelium in the soil is prevented by enclosing each seed in a narrow glass cylinder (6 × 2 cm.) which is pressed into the soil. Counts of diseased seedlings are made at stated intervals.

The *drilling capacity* of disinfected grain is determined by the use of an experimental drilling machine. Three estimations are made of the amount of grain which passes through the machine after it has made one hundred revolutions.

Field experiments are carried out in addition to laboratory tests.

The *hygroscopic nature* of dust preparations is determined by exposing a known quantity of the dust in a moisture-saturated chamber. The water increase by weight is estimated after a period of seven days.

Disinfectant dusts are tested for their *action upon iron*. Layers of the dust are placed upon small iron plates, some of which are then put in a moisture-saturated chamber and others in the laboratory, the moisture of the air of which is determined. The rust-promoting properties of the disinfectant are assessed after a period of three days.

THE MAIN TESTS

Materials which successfully pass the preliminary tests are recommended for the main tests, the chief object of which is to determine the efficacy of the preparation in practice under different weather and soil conditions. The main tests are carried out by at least eight and not more than twelve of the central stations for plant protection, and are made in much the same way as the preliminary tests. Only those concentrations which have proved to be effective in the preliminary tests are used in the main tests.

After the completion of the main tests the results are considered by a judging committee which decides whether the material shall be included in the official list. For materials giving unsatisfactory results and not admitting of a definite decision, the manufacturer is given an opportunity of furnishing his observations before a meeting of the committee.

When a material has been passed for inclusion in the official list, the manufacturer is required to sign an agreement which tends to safeguard the purchaser and among other things guarantees that no other products will be placed on the market under the trade-mark of the listed preparation.

So far as I am aware, the submission of material for official tests is on a purely voluntary basis and this seems to me to be rather a strong point in favour of such a scheme. Nobody is precluded from the manufacture and marketing of plant protective substances. Those who do so have the opportunity of submitting their products for official tests which, if successfully passed, allow it to be known that the material is officially approved for use at certain concentrations by certain methods for certain purposes of plant protection. This should mean a great deal if the work of official stations enjoys the confidence of the farming community.

Apart from the official work carried out at the State Biological Institute, work on an elaborate scale is conducted by the large industrial concerns engaged in the manufacture of plant protective substances. The Biological Institutes of the I.G. Farbenindustrie at Leverkusen (Cologne) and Hoechst (Frankfurt) have large well-equipped buildings with adequate field facilities close at hand. The testing work carried out by the scientific staffs of these Institutes closely approaches the lines followed by the Official Institute and results obtained by the station at Leverkusen are checked at Hoechst. I believe that many thousands of materials were tested before the present preparation recommended by I.G. Farbenindustrie for seed disinfection was marketed. While visiting these institutes I had the privilege of seeing tests being made with preparations in order to determine their efficiency for controlling the following diseases: Bunt

in wheat, Leaf Stripe in barley, Smut in oats, *Fusarium* disease of rye, Apple Scab and Downy Mildew of the vine. An entire laboratory was devoted to testing work dealing with fungicides for the prevention of timber decay. Apart from fungicides, insecticides and vermicides were receiving equally careful attention. From the few observations I was able to make, the relations existing between the staffs of the industrial institutes and that of the Official Institute, seemed to be healthy and of the happiest kind.

During the subsequent discussion Mr C. T. Gimingham referred to a general point touched on by Mr Marsh, and emphasized the importance of keeping in mind the precise object in view in working out the technique of laboratory methods of testing.

Methods may be required (*a*) for use in fundamental research on toxicity and (*b*) in testing commercial and other preparations. For the first purpose it is justifiable and necessary to aim at the highest possible degree of accuracy, but to attain this in biological test methods almost invariably involves a degree of elaboration and expenditure of time which is likely to disqualify the method for use in routine work. In devising methods for the second purpose it may be necessary to sacrifice some degree of refinement, and it will be agreed that it is likely to be waste of effort to use elaborate and very accurate apparatus unless it is also possible to have standardized and uniform biological material for the tests. In this connexion Mr Gimingham referred briefly to the methods adopted by the German Plant Protection Service for use in their official tests, some of which he had had an opportunity of seeing during a visit to the Institute at Dahlem last summer. He had formed the impression that at any rate so far as the methods for testing insecticides were concerned (and probably the same applied to the methods for fungicides) they had been designed with a very practical aim. Although giving satisfactory and reliable results for the particular purpose in view they would not all necessarily be suitable for (nor are they intended for) more fundamental investigations. Relative simplicity and ease of operation must not be lost sight of in considering methods for routine testing of commercial products.

SOME CHYTRIDIACEOUS FUNGI FROM NORTH AFRICA AND BORNEO

BY F. K. SPARROW, JR.

(With 2 Text-figures)

SINCE our knowledge of the distribution of Chytridiaceous fungi is so fragmentary, it would seem of interest to record in this brief note the occurrence of certain of these organisms in two rather unusual geographic localities.

The material from North Africa appeared in gross water cultures started from a bit of vegetable "trash" collected in a small pond near Tangier in April 1933, by Mr E. F. Warburg, M.A., of Trinity College, Cambridge. The specimens from Borneo, excellently preserved in formalin, were found in a collection of fresh-water red algae made by Mr P. W. Richards, M.A., also of Trinity College. To these two gentlemen the writer wishes to express his best thanks.

TANGIER

In the cultures started from a bit of moist vegetable trash, there appeared after a few days a profuse growth of *Chlamydomonas* spp. and *Protoderma* sp. While several Chytridiaceous fungi were found on these two algae, in only one were significant phases in the life history observed. This fungus, however, proved to be of very great interest, and a comparison with other related forms indicated that it belonged in a genus of its own which was possibly allied to *Sporophlyctis*. The name *Sporophlyctidium africanum* has been given to it in a previous paper⁽¹⁾.

The fungus has an exceedingly simple structure and mode of development. The zoospore, lying free in the water, produces a highly refractive, wedge-shaped germ tube (Fig. 1a, b). The tip of this structure penetrates the cell wall of the alga. No further development within the host cell takes place. Subsequent growth results in a differentiation of the germ tube and the body of the spore, the former becoming more isodiametric and stalk-like, the latter larger and obpyriform (Fig. 1d). Although no cross-wall could be detected separating these two portions of the thallus, when the fungus had attained its full size the granular content of the more distal portion was in marked contrast to the refractive material within the stalk. Ultimately, the content of the distended part became divided into four to

six spores which were liberated through a small lateral pore formed in the wall. Outside, they remained grouped for a time before the orifice but ultimately this cluster fell apart and the spores floated away. Each was spherical, 2μ in diameter, possessed a minute oil globule and was devoid of cilia. The latter point was ascertained after a close study of a number of spores from various sporangia. After discharge the sporangium and its stalk collapsed (Fig. 1c, d).

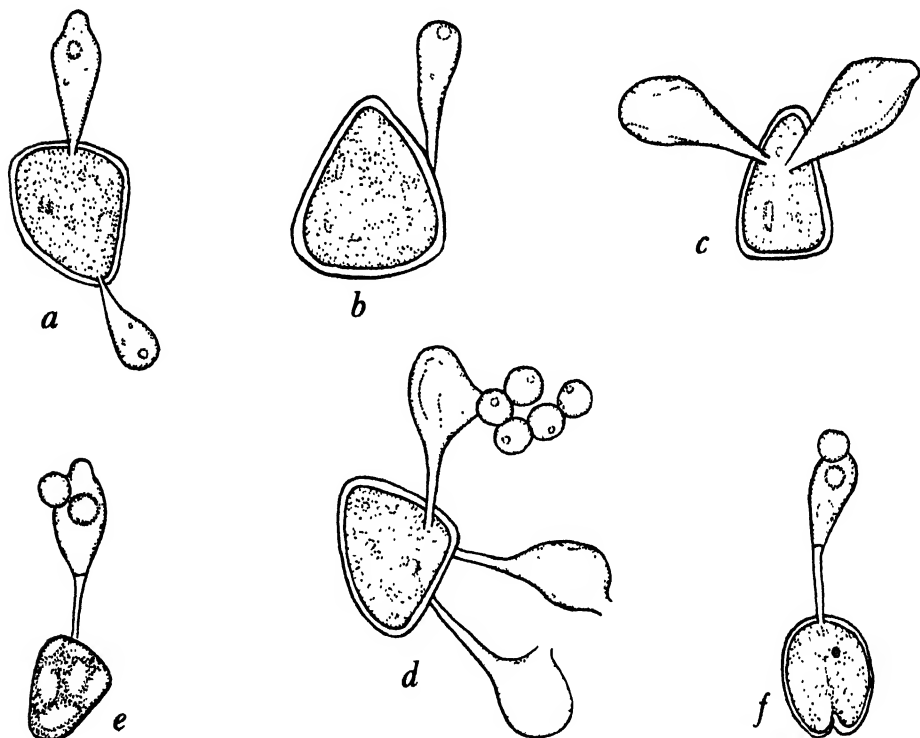


Fig. 1. *Sporophlyctidium africanum*, parasitic on *Protoderma* sp., Tangier. a-d, $\times 2500$; e, f, $\times 1500$. a, b, stages in the formation of the sporangia; c, empty sporangia; d, discharging sporangium (upper), two empty sporangia below; e, f, possible resting spores with companion cells.

What may possibly be the resting spores of this fungus were observed in a few instances several weeks after the sporangial stage had been found. These were similar in shape to the sporangia, 7μ long by 4μ in diameter, and possessed a somewhat thickened wall and a single large oil globule in the content (Fig. 1e, f). Attached laterally to the upper portion of the spore was an empty, spherical cyst which undoubtedly functioned as a "male" or "companion" cell. Germination of these spores was not observed, nor were early stages of the sexual process seen.

In its main features this interesting organism closely resembles *Sporophlyctis rostrata* Serbinow, a form found parasitic on *Draparnaldia* in Russia⁽²⁾. In the latter fungus the relationship of the parasite to the alga is similar to that of the present form, only the tips of the rhizoids penetrating the host cell. In the shape of the sporangium, position of the discharge pore, and the non-ciliation of the zoospore this resemblance is further emphasized. However, in certain essential features, the African fungus differs markedly from *Sporophlyctis*. In the former, the structure which makes contact with the host cell is never branched or rhizoidal in character, but unbranched and "inflated" (broad). Nor are the zoospores of this fungus formed in a vesicle extruded from the sporangium, but rather are fully formed within the latter body, and discharged individually. Further differences between the two are found in the sexual stage. In *Sporophlyctis* there is a copulation between two thalli, a large, male plant similar to the sporangium and a smaller, narrowly clavate, spiny-walled female one. A portion or all of the content of the larger thallus passes over into the smaller one and the latter is ultimately transformed into a resting spore. While early stages in the formation of the resting spore of the African fungus were not observed, it is apparent that the receptive individual is the larger of the two; also, the resting spore thus formed is smooth-walled. No evidences of any rhizoidal development were found on this male cell, although such may have been present in the early stages of conjugation.

These differences between the African fungus and *Sporophlyctis* seemed of such a fundamental nature that a new genus was proposed for its disposition. This has been given, in Latin, in a previous paper⁽¹⁾. For convenience, the following English generic and specific diagnoses are appended:

Sporophlyctidium. Zoospore lying free in the water, upon germination producing a single, unbranched, somewhat inflated tube, the tip of which penetrates the host cell; the body of the zoospore expanding and becoming the sporangium. Zoospores formed in the sporangium, at maturity liberated from the latter body after the deliquescence of a papilla; motionless, without cilia. Resting spore extramatrical, with a companion cell.

S. africanum. Sporangium narrowly pyriform, smooth-walled, colourless, the narrower end continuous with the unbranched germ tube; $5\ \mu$ long by $3.5\text{--}4\ \mu$ in diameter; forming a single subapical pore through which the spherical, uniguttulate, non-ciliated spores, $2\ \mu$ in diameter, are extruded. Resting spores similar in shape to sporangia, $7\ \mu$ long by $4\ \mu$ in diameter, with a spherical companion cell $3\ \mu$ in diameter; germination not observed.

Parasitic on *Protoderma* sp., Tangier, North Africa; collected by E. F. Warburg, April 1933.

Three other Chytridiaceous fungi were found in the same material that yielded *Sporophlyctidium*. The first of these (Fig. 2 a), also found on *Protoderma*, possessed broadly pyriform, extramatrical sporangia, $9\ \mu$

high by 7μ in diameter. At the apex were two broad, blunt papillae, 2.5μ high by 3.5μ in diameter. Within the algal cell a short peg was noted but whether or not it was branched could not be determined. No further stages were observed.

The other two fungi were found on resting cells of *Chlamydomonas* sp. One (Fig. 2c) possessed extramatrical, somewhat angular resting spores about 12μ in greatest diameter. The spores were thick-walled and their surface provided with irregularly disposed, blunt, solid bullations. Where the host content was contracted, a branched rhizoidal system could sometimes be detected attached to these resting spores. The sporangial stage was not observed. The other fungus found on *Chlamydomonas* (Fig. 2b) had smooth, thick-walled, spherical, extramatrical resting spores, $6-9\mu$ in diameter, each of which possessed a spherical, empty companion cell, $2.5-3\mu$ in diameter. Parts of a branched rhizoidal system could occasionally be detected within the alga. No sporangial stage was found.

These three fungi are probably all species of *Rhizophidium* and referable to European species, but until more is known of their life history, they cannot be identified.

BORNEO

Four Chytridiaceous fungi were found in the material, preserved in formalin, brought back by Mr Richards.

In material of green algae from a drainage ditch at the edge of the forest near Claudtown, Sarawak, collected in December 1932, many specimens of a *Synedra* which were infected with *Rhizophidium fusus* Zopf were found. The sporangia (Fig. 2d) were fusiform, $10-15\mu$ long by $4-6\mu$ at their greatest diameter, and were nearly always slightly tilted. The intramatrical system consisted of a series of delicate, branched rhizoids. These specimens differed in no significant respect from the common European or American ones.

In material of *Cosmarium* sp. from this same locality there were found many discharged sporangia of a very small species of *Rhizophidium* (Fig. 2e) which appeared to be closely allied to *R. carpophilum* Zopf, or *R. minutum* Atk. The nearly spherical sporangia were $5-7\mu$ in diameter, with a relatively broad apical pore. The rhizoidal system was delicate and sparingly branched. In a few undischarged mature

Legend for Fig. 2.

Fig. 2. a, sporangium of *Rhizophidium* (?) on *Protoderma*, Tangier, $\times 2500$; b, resting spores of *Rhizophidium* sp. (?), with companion cells, on *Chlamydomonas*, Tangier, $\times 2500$; c, resting spore of *Rhizophidium* sp. (?), on *Chlamydomonas*, Tangier, $\times 1300$; d, *Rhizophidium fusus* on *Synedra*, Borneo, $\times 1300$; e, *Rhizophidium* sp. (?) on *Cosmarium* Borneo, $\times 1300$; f-h, sporangia of *Rhizophlyctis borneensis* n.sp., on diatoms (diat.), Borneo, $\times 1300$; i, *Chytridium Schenkii* (?), on *Cosmarium* sp., Borneo, $\times 1300$.

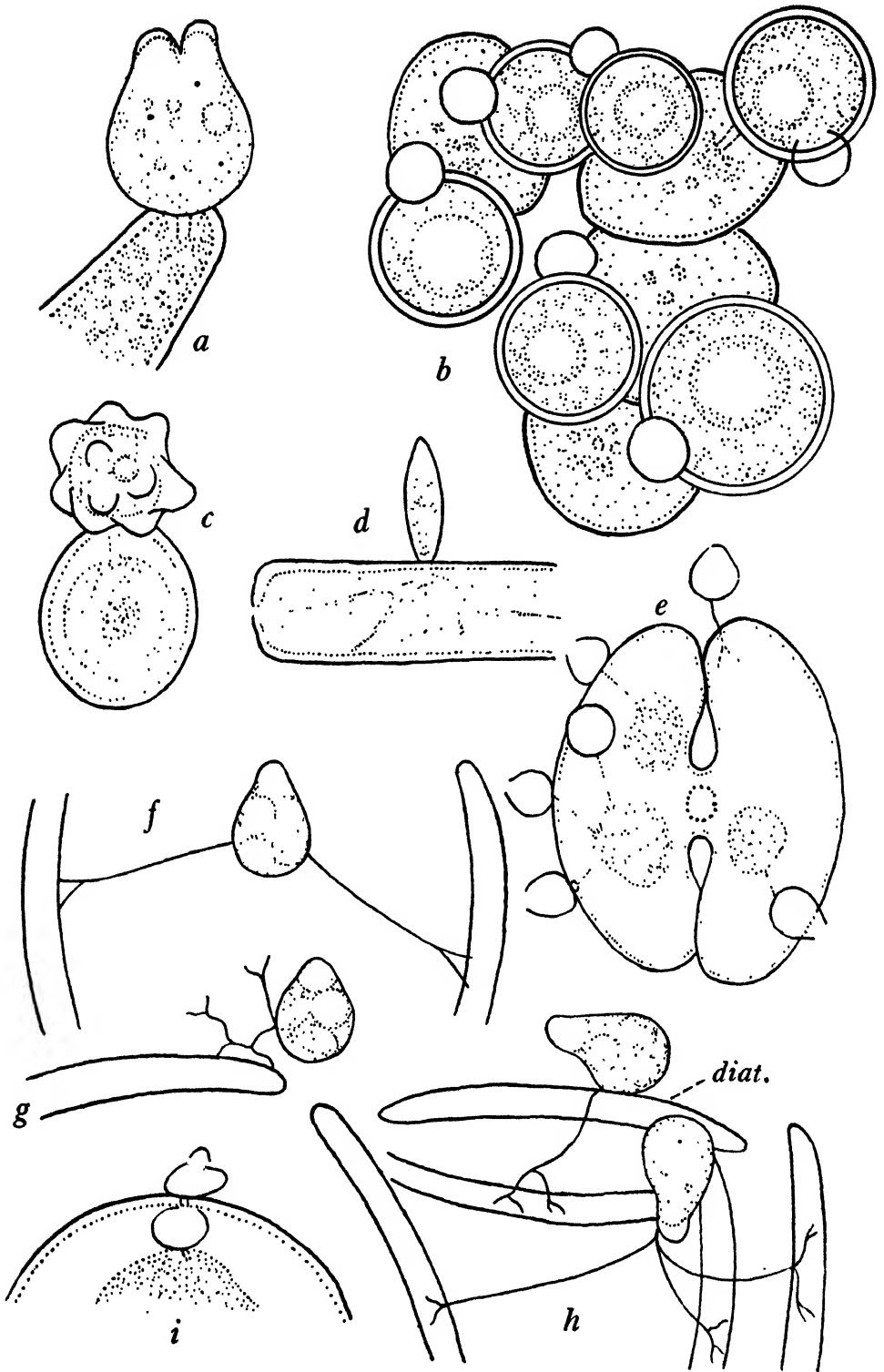


Fig. 2. (See opposite for legend.)

sporangia, eight to ten regularly placed oil globules were seen. Each of these globules probably marked the position of a zoospore within the sporangium.

On *Cosmarium* from the same locality several very small specimens of a *Chytridium* were found. The sporangia, which were only about 8μ long by 5μ in diameter, were somewhat elliptical, inclined at an angle to the wall of the host, and possessed a subapical pore. Attached in the vicinity of this pore was a strongly convex operculum (Fig. 2i). Within the host cell there was a spherical subsporangial swelling about 5μ in diameter from which several rhizoids arose. In its general aspect this Chytrid resembles *Chytridium Schenkii* although the small size of the parts and the subapical position of the exit pore mark it off from this species.

Among diatoms epiphytic on a fresh-water Floridean (No. 2585), collected in a small forest stream near Long Kapa, Mt Dulit, Ulu Tinjar, Sarawak in November 1932, there were found a number of plants of a small species of *Rhizophlyctis*.

The sporangia (Fig. 2f-h) were pyriform or bursiform, $12-15\mu$ long by $7-10\mu$ in diameter, with a prominent, broad, blunt papilla. From the middle or lower part of the sporangium there were produced one to three very delicate rhizoids, the tips of which made contact with different diatom frustules. Generally these rhizoids remained unbranched until reaching the vicinity of the diatom where they branched profusely.

Within many sporangia four to six somewhat spherical zoospores, about 4μ in diameter, could be observed. On several of these aguttulate spores, which were found at the orifice of a partially discharged sporangium, a single cilium could be detected.

In the shape of its sporangium this fungus closely resembles *Rhizophlyctis mastigotrichis* (Nowakows.) Fischer. However, it has much smaller sporangia ($12-15 \times 7-10\mu$, cf. 40μ diameter), the rhizoids are more delicate and attenuated, not swollen distally, and a smaller number of zoospores (four to six, cf. fifty or more) are produced in each sporangium.

Because of these well-marked differences, which are apparent even in preserved material, I consider the fungus distinct from *R. mastigotrichis*, its closest fresh-water relative, and name it *Rhizophlyctis borneensis*.

R. borneensis n.sp. Sporangium free, pyriform or bursiform, $12-15\mu$ long by $7-10\mu$ in diameter, smooth-walled, with a broad papilla; rhizoids one to three, arising from the middle or lower part of the sporangium, generally branching in the vicinity of the host cell, polyphagous. Zoospores spherical uniciliate, four to six formed in a sporangium, 4μ in diameter. Resting spore not observed. Parasitic on diatoms, British North Borneo. Coll. P. W. Richards.

RHIZOPHLYCTIS BORNEENSIS sp.nov.

Sporangiis liberis, piri- vel bursiformibus, 12–15 μ longis, diametro 7–10 μ , membrana levi cinctis, papilla lata ornatis; *rhizoidibus* 1–3, sporangio medio vel parte inferiori productis, juxta matricem fere ramosis, polyphagis; *zoosporis* sphaericis, uniciliatis, in sporangio 4–6, diametro 4 μ .

Hab. ad Diatomaceas in Borneo Brit. Septentr.

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A RECORD OF TWO YEARS' CONTINUOUS OBSERVATIONS ON *BLASTOCLADIA* *PRINGSHEIMII* REINSCH

By DAPHNE LLOYD, B.Sc.

Royal Holloway College

(With 3 Text-figures)

THE aquatic fungus, *Blastocladia Pringsheimii*, was first described in Germany by Reinsch⁽¹⁷⁾ in 1878. Since then it has been described in Germany by von Minden^(13, 14), in the United States of America by Thaxter⁽²²⁾, Kanouse^(8, 9), Cotner⁽⁴⁾, and Sparrow⁽¹⁸⁾, and in Denmark by Petersen^(15, 16). It was first recorded in Great Britain in 1930 by Barnes & Melville⁽¹⁾; and it has been reported three times more recently, in 1933-4 by Cooke & Forbes⁽³⁾, in 1935 by Forbes⁽⁵⁾ and in 1936 by Sparrow⁽²¹⁾.

B. Pringsheimii has not, however, been described in detail in Britain, and its life history is not yet completely known. It seemed worth while therefore to make a further investigation of it. A special opportunity offered itself when the fungus was found growing abundantly under natural conditions in three pools and a stream in the grounds of Royal Holloway College. Although *B. Pringsheimii* is apparently saprophytic and will grow on various twigs and fruits in water, it has not yet been successfully cultured in artificial media.

SOURCE OF MATERIAL

Attempts were made to cultivate the fungus in the laboratory in nutrient solutions and on nutrient agar media, but it was found impossible to maintain normal growth for any length of time. Plants were transferred from the pond to a solution of soil extract containing a 0.5 % solution of potassium nitrate and to a 1 % pea solution, and they were mounted in their own pond water on various agar media. Zoospores swarmed and germinated in these solutions and on these media, but, by the time the first sporangium was initiated, or earlier, growth ceased and the young plants showed an unhealthy accumulation of oil. Agar films containing and supported by a mesh of cotton-wool were suspended in the ponds to catch the zoospores, but these proved useless, as the agar slowly dissolved in the water before there

was any inoculation. Since none of these three methods proved very successful, the investigation was confined almost entirely to material brought from a pond in the botanical greenhouse. For two or three hours after collection normal growth appeared to continue, and the observations on this material were perhaps even more reliable than those made under the best of artificial conditions, as is evident from the later account of the swarming of the zoospores.

METHOD OF INVESTIGATION

The material was collected by von Minden's⁽¹⁴⁾ method of suspending fruits in a wire cage six to twelve inches below the surface of the water. Tomatoes were finally selected as the best fruit, as their thin transparent skin renders observation of distribution of individual thalli easier, and the soft pulp separates relatively easily on dissection from the rhizoids of the fungus. The tomatoes were suspended for a period of three to fourteen days in the water of the greenhouse pond, of which the temperature varies between 13° and 18° C., when the skin became covered with white pustules each containing from one to forty independently established plants. The tomatoes were then washed in running tap water in the laboratory and replaced in water from the greenhouse pond and examined immediately.

Since material was abundant, culture in the laboratory was not essential, but it was useful for comparison. When it was found that sporangia failed to discharge zoospores in culture in either pond water or sterile distilled water, that the water became cloudy with bacterial zooglea after the tomatoes had been kept in the laboratory for one or two days, and that inoculations of fresh tomatoes in sterile distilled water produced immature plants which developed only rudiments of sporangia, such cultures were discontinued, and aquaria were made with pond mud, pond water, green water plants, snails, etc. In these surroundings *B. Pringsheimii* continued to develop normally in the laboratory, and fresh sterile tomatoes were inoculated and produced normal healthy individuals.

For examination, two or three living pustules were mounted together in a hanging drop or in a moist chamber. The latter was generally used, because more individuals could be mounted together. Zoospores were discharged more readily in pond water, or in a solution of soil extract containing potassium nitrate, than in distilled water. Early stages of germination of zoospores were observed in pond water, while young plants have been obtained in a hanging drop of pond water on an agar film.

Complete plants were fixed and stained for the study of the development of the sporangium and of the zoospores within the sporangium. Since the wall of the sporangium is thin and permeable and the

diameter of the sporangium is small, sections are not essential provided that de-staining is adequate. Several fixatives were tried, and Nawaschin's fixing fluid was found most satisfactory. The material was well washed after fixation and stained by Newton's gentian violet method (La Cour⁽¹²⁾). The plant was mounted under a cover-slip in 0.1 % water solution of gentian violet for five minutes, and the cover-slip tapped gently, so that the sporangia spread out like a fan from the main axis.

Sections of plants were made for studying finer detail. Whole pustules were fixed, dehydrated and embedded in paraffin wax, or single plants were embedded after fixation in agar films, and the small blocks of agar were then embedded in paraffin wax. By the latter method approximately longitudinal or transverse sections of the main axis and sporangia could be more readily obtained. Sections were cut 6–10 μ thick and stained as above.

OBSERVATIONS

Form of thallus and position of sporangia on thallus

The thallus is differentiated into a rhizoidal system, which branches extensively as it penetrates into the substratum, and an emergent, sporangium-bearing axis, which exhibits such wide variation in its growth form that sometimes a second species, *B. globosa*, has been suspected. Thin-walled sporangia are developed at the tips of the usually branched main axis; they are cylindrical but vary considerably in length (300–110 μ) and breadth (30–15 μ). In addition, there may be produced simple and branched filamentous hairs and resting spores, the latter spherical, ovoid or shortly cylindrical with a wall of three layers, the middle layer being perforated.

Records of the variation in growth form and in frequency of resting spores and hairs have been kept for seventeen months, and these appear in Tables I and II. Table I gives the observations on material from the greenhouse pond, where the temperature varied between 13° and 18.5° C., and Table II gives observations on material collected from a pool in the botanical garden, with a temperature varying between 0° and 17.5° C. Each entry represents the range of variation within six pustules. Since the date of inoculation, and hence the true age of the plant, was not known, the maximum age or greatest possible age, that is, the time of immersion of the material, is given. The differences in growth form were referred to four types A, B, C and D; type A includes forms with a globose main axis; type B, forms with a sturdy main axis with a swollen or lobed apex; type C, forms with a sturdy branched main axis, and type D, forms with a slender, branched, filamentous main axis. Representative plants from these four classes are figured (Fig. 1).

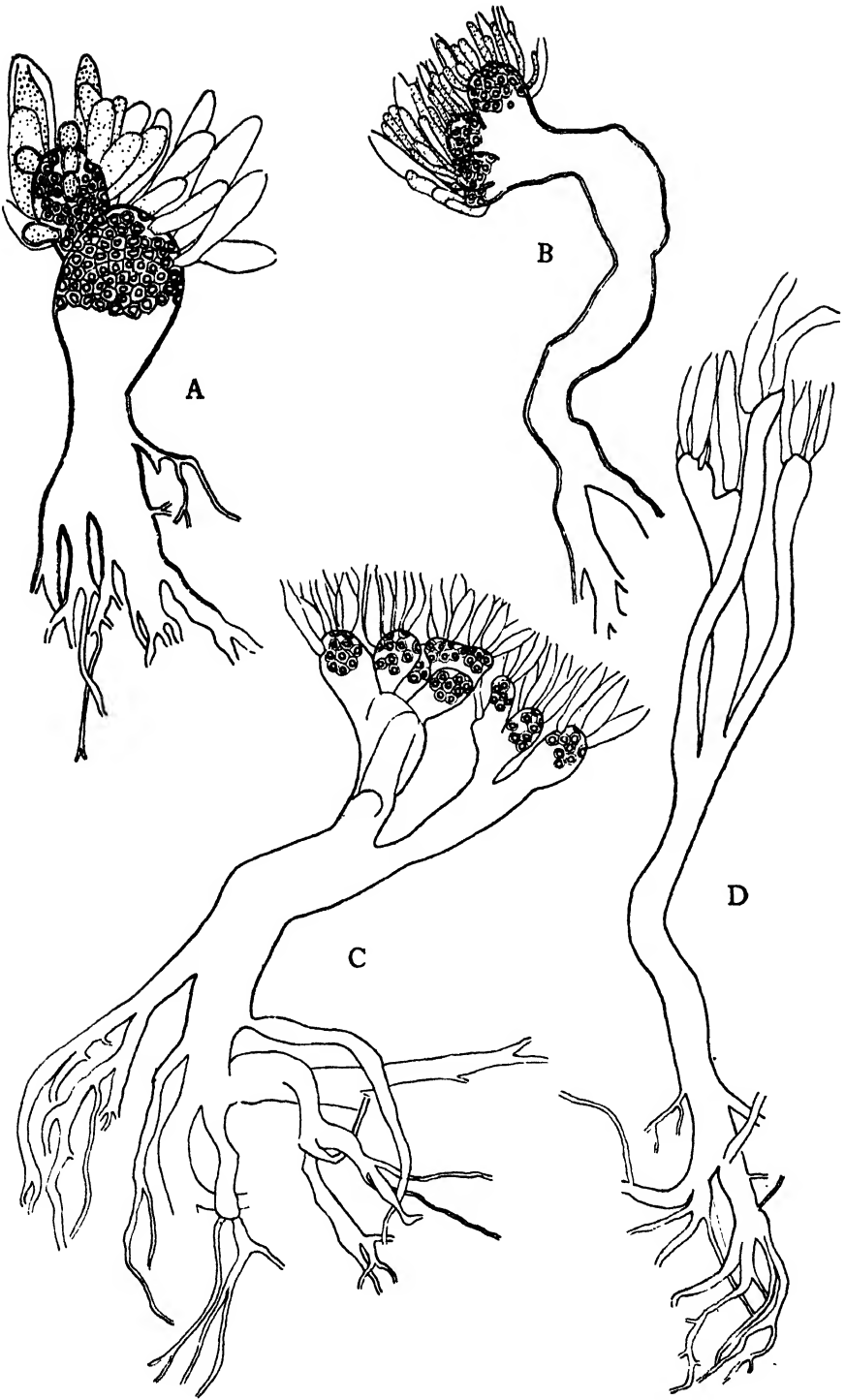


Fig. 1. Four plants ($\times 80$) of *Blastocladia Pringsheimii* Reinsch, showing variation in form and referred to, in the text, as types A, B, C, D. All bear empty sporangia. A and B have young sporangia. A has five resting spores. A, B and C show scars where sporangia have fallen.

A scrutiny of Tables I and II suggests that there is no periodicity in growth form or in frequency of resting spores and hairs. The forms of thallus included under B and C were most common in both pools, and the four types were developed at all seasons. Although there was

Table I

Date	Maximum age in days	Growth form (see Fig. 1)	Resting spores, frequency: F, frequent; O, occasional; A, absent	Hairs	
				Frequency	Type: S, simple; B, branched
1. v. 34	60	A, C	F	O	S
15. v. 34	3	A	A	O	S
17. v. 34	24	A, C	F	A	
27. v. 34	7	B, C	A	O	S
4. vi. 34	10	B, C	O	O	B
15. vi. 34	6	B, C	O	O	S
17. vii. 34	14	A, B, D	F	O	S & B
28. vii. 34	11	B, C, D	F	A	
4. viii. 34	7	D	F	A	
11. viii. 34	7	A, B, C	A	O	B
16. viii. 34	5	No inoculation			
27. viii. 34	11	B, C	O	O	S
3. ix. 34	7	A, C	A	A	
10. ix. 34	7	C, D	A	A	
17. ix. 34	7	B, C	A	A	
24. ix. 34	7	B, C	A	A	
1. x. 34	7	B, C, D	A	F	S
13. x. 34	12	B, C	A	O	S
26. x. 34	21	B	A	A	
30. x. 34	11	C	O	A	
13. xi. 34	10	B, C	A	O	B
29. xi. 34	10	B	A	A	
14. xii. 34	17	C	O	O	B
27. xii. 34	13	B, C, D	F	A	
7. i. 35	11	B, C	A	O	S & B
24. i. 35	17	C	O	A	
13. ii. 35	13	B, C	A	O	B
21. ii. 35	10	A, C	F	A	
6. iii. 35	12	C, D	O	A	
30. iii. 35	12	B, C	F	A	
8. iv. 35	10	B, C	A	A	
18. iv. 35	10	B, C	O	O	S & B
7. v. 35	10	B, C, D	F	O	S
24. v. 35	13	C	A	O	S & B
14. vi. 35	21	B, C	F	A	
27. vi. 35	14	C, D	F	O	B
12. vii. 35	14	C, D	F	A	
26. vii. 35	14	B, C, D	F	A	
9. viii. 35	14	A, C, D	F	F	S
11. ix. 35	33	A, B, C	F	A	

no seasonal periodicity in the development of resting spores, they were found most abundantly in the summer months; during the winter they appeared occasionally in the greenhouse pool and very occasionally indeed in the garden pool. The filamentous hairs were likewise not always present, and the occurrence of both branched and

unbranched hairs was related neither to the presence of resting spores nor to the time of year.

Since the globose type of thallus (A) occurred irregularly in these records, and as measurements of sporangia on these plants came within the range which has been described for *B. Pringsheimii*, it

Table II

Date	Maximum age in days	Growth form (see Fig. 1)	Resting spores, frequency	Hairs	
				Frequency	Type
1. v. 34	60	B, C	A	O	S
8. vi. 34	42	A, B, D	F	O	S
8. vi. 34	14	B	A	F	S
17. vii. 34	14	A, C	F	A	
28. vii. 34	11	A, C	A	O	S & B
4. viii. 34	7	B, C	F	A	
11. viii. 34	7	A, B	A	A	
16. viii. 34	5	A, B	A	O	S
27. viii. 34	11	B	A	A	
3. ix. 34	7	C	A	A	
10. ix. 34	7	C	A	O	B
17. ix. 34	7	C	A	A	
24. ix. 34	7	C, D	A	A	
1. x. 34	7	B	A	A	
13. x. 34	12	B	A	O	S
26. x. 34	21	B, C	O	A	
13. xi. 34	10	No inoculation			
29. xi. 34	10	D	A	A	
14. xii. 34	17	B, C	O	A	
27. xii. 34	13	B	A	O	S & B
7. i. 35	11	No inoculation			
5. ii. 35	19	C	A	A	
21. ii. 35	10	B, C, D	A	A	
21. ii. 35	21	B, C, D	A	A	
6. iii. 35	56	B, C	A	A	
30. iii. 35	14	C, D	A	O	B
8. iv. 35	10	B, C	A	A	
18. iv. 35	10	C, D	A	A	
7. v. 35	10	B, C	A	O	S & B
16. v. 35	9	C	A	F	S & B
12. vi. 35	14	B, C	F	A	
28. vi. 35	14	B, C	F	A	
12. vii. 35	14	A, C	F	A	
26. vii. 35	14	A, C	F	A	
9. viii. 35	14	A, C	O	O	S
11. ix. 35	33	C	F	A	

appears that the globose thalli are growth variations of *B. Pringsheimii* and not those of a separate species. All grades of variation between the globose type A and the more slender branched type D were found. A single pustule often contained plants which could be referred to two or more of these types. These variations in growth form appear, therefore, to belong to a single species.

Development of sporangium and zoospore

The observations on the development of the sporangium and zoospore made on living material have been amplified by a more intensive study of fixed and stained material. The sporangium arises as a small papilla which is very soon cut off by a wall from the main axis. It may then contain five to twenty nuclei, and the not infrequent absence of nuclei in the main axis immediately below the young sporangium suggests that it has been cut off by the wall after these few nuclei have passed in. Simultaneous divisions of these nuclei follow, and since the sporangium finally contains 150–250 nuclei, the nuclei must divide several times in the development of the sporangium. Dividing nuclei have been observed. It may be calculated that the formation of the final number from five nuclei would involve five simultaneous nuclear divisions. The determination of the frequency of these divisions has presented difficulties, as it has not been possible to follow the complete development of an individual sporangium in culture. There is evidently no diurnal periodicity in nuclear division, for material fixed at different times of day shows similar nuclear figures, and all stages of development of sporangia are seen at any one time. It is more probable that nuclear division occurs at definite intervals after initiation of the sporangium, and so is related to the stage of development. The frequency of nuclear division has been surmised thus: tomatoes, which had been immersed only two days in the pond, showed plants with papillae, while after four days' immersion they bore sporangia. These sporangia had developed from the papilla stage in two days at the most, and the nuclei must have divided at least twice a day. This simultaneous nuclear division is not confined to the sporangium alone; the main axis shows similar regularity in nuclear division.

After the last division has taken place in the sporangium, the nuclei go into a resting condition, and only a central nucleolus surrounded by a colourless region can be distinguished. At the same time the cytoplasm begins to divide up by cleavage vacuoles. Finally, when the protoplasm has been cut up into zoospores, there is a large proportion of deeply staining material in each zoospore. The large "sub-triangular" nuclei of Thaxter are readily discerned within the sporangium immediately before emergence.

*The necessary conditions for dehiscence of the sporangium
and emission of zoospores*

It has frequently been said that the zoospores do not always emerge from the sporangium, and the necessary conditions for emergence are not yet fully understood. Thaxter⁽²²⁾ who first described the zoospores of *B. Pringsheimii* says: "As the fungus develops, growing as it almost

invariably does in tufts, it forms the centre of a dense mass of bacteria, which finally choke the sporangia completely; so that as a rule only those first formed are able to discharge their contents. As a result the zoospores commonly die without escaping." Petersen⁽¹⁶⁾ made a similar observation. Kanouse⁽⁹⁾ found that "zoospores swarm abundantly and readily in cultures that are two weeks old or more after the plants are transferred to distilled water and the temperature of the room in which the study is being pursued is considerably below normal". Cotner⁽⁴⁾ showed that above 14° C. and below 11° C. there is incomplete cleavage of the protoplasm resulting in the formation of abnormal spores or absence of swarming, and thus explains the earlier observations.

Since old plants from the ponds and laboratory aquaria of Royal Holloway College showed, with rare exception, only empty sporangia, it must be concluded that the sporangia discharge freely under natural conditions. It is clear that, whereas the conditions existing during *emission* are not so important, those under which the *formation* of zoospores take place are very important. Swarming, when it occurred, was usually observed soon after the material had been brought into the laboratory. The sporangia generally ceased to discharge zoospores after the plant had been mounted for three hours, but exceptionally, the emergence of zoospores was observed in a hanging drop after eight, thirteen or twenty-two hours. Sporangia discharged zoospores even after the main axis had been torn in dissection, if the sporangia were in a mature state when the material was brought into the laboratory. Yet the conditions existing during emission must have some effect, because mature sporangia in fresh material did not always discharge their spores.¹

Observations on living material were made at least once a week during the six terms of the sessions 1933-5. Fields of zoospores have been watched, and the zoospores from fifty-three sporangia were followed from emergence to germination. The temperature at which zoospores emerged was between 15° and 18° C. This is rather higher than the optimum temperature for zoospore formation given by Cotner, but the pond from which the plants were collected was at a correspondingly higher temperature (13°-18° C.) than his source of material, and this may account for the difference. The results are given in Table III and may be analysed as follows:

(1) Zoospores were emitted at all seasons of the year and at all times of day.

¹ Variations in temperature, light intensity and acidity of the medium probably influence the emergence of the zoospores, because these factors affect their later development. Zoospores do not germinate after prolonged exposure to artificial light, and the greatest development of sporelings was obtained on a medium with a pH value the same as that of the pond water, i.e. pH 8.0.

Table III

Date	Time of dehiscence of sporangium	Number of discharging sporangia observed	Germination of swimmers observed	Occurrence of resting spores on the same plant
12. xi. 33	6.45 p.m.	1	+	—
12. xi. 33	8.30 p.m.	2	+	—
13. xi. 33	7.20 a.m.	1	+	—
13. xi. 33	9.30 a.m.	2	+	—
13. xi. 33	...	2	+	—
29. xi. 33	10.45 a.m.	1
2. xii. 33	8.50 a.m.	1	+	—
2. xii. 33	11.5 a.m.	2	+	—
16. ii. 34	...	1	+	...
13. iii. 34	5.45 p.m.	Several	+	...
14. iii. 34	10.0 a.m.	Several	+	...
15. iii. 34	9.30 a.m.	Several	+	...
28. iv. 34	11.0 a.m.	1	+	...
3. v. 34	10.15 a.m.	1	+	...
5. v. 34	10.45 a.m.	1	+	...
7. v. 34	11.30 a.m.	1	+	...
2. vi. 34	10.45 a.m.	1	+	...
9. vi. 34	...	Several
23. x. 34	5.15 p.m.	3	...	—
23. x. 34	5.30 p.m.	2	...	—
20. xi. 34	9.45 a.m.	1	...	—
20. xii. 34	10.2 a.m.	2	...	—
5. xii. 34	5.15 p.m.	1	0	—
6. xii. 34	7.45 a.m.	1	+	—
6. ii. 35	9.55 a.m.	1	0	...
6. ii. 35	10.0 a.m.	2	0	...
6. ii. 35	10.2 a.m.	3	0	...
6. ii. 35	10.14 a.m.	4	0	...
6. ii. 35	10.52 a.m.	5	0	...
6. ii. 35	10.55 a.m.	6	0	...
6. ii. 35	10.57 a.m.	7	0	...
6. ii. 35	10.57 a.m.	8	0	...
20. ii. 35	9.30 a.m.	1	+	—
20. ii. 35	9.55 a.m.	2	+	—
20. ii. 35	10.0 a.m.	3	+	—
20. ii. 35	10.10 a.m.	4	+	—
20. ii. 35	10.50 a.m.	5	+	—
20. ii. 35	10.55 a.m.	6	+	—
13. iii. 35	10.15 a.m.	Several	0	—
8. v. 35	9.30 a.m.	1	0	—
8. v. 35	10.0 a.m.	2	0	+
8. v. 35	10.15 a.m.	3	0	+
8. v. 35	10.15 a.m.	4	0	+
9. v. 35	9.30 a.m.	1	0	+
9. v. 35	11.30 a.m.	2	+	+
12. vi. 35	10.15 a.m.	1	+	—
12. vi. 35	10.50 a.m.	2	+	—
12. vi. 35	10.55 a.m.	3	+	—
12. vi. 35	11.0 a.m.	4	+	—
12. vi. 35	11.40 a.m.	5	+	—
12. vi. 35	11.40 a.m.	6	+	—
12. vi. 35	11.50 a.m.	7	+	+
12. vi. 35	12.15 a.m.	8	+	—

(2) Except in eleven of the fifty-three instances, two or more zoosporangia discharged zoospores simultaneously or nearly so. There was therefore ample opportunity to observe fusion of gametes, if such they were.

(3) The zoospores usually settled down and germinated within two hours. Where no germination is recorded in Table III the zoospores had been watched for two to three hours by artificial light at a high temperature (sometimes as high as 37° C.).

(4) The sporangia, which discharged zoospores, were generally borne on plants which did not produce resting spores. But where resting spores were present, the zoospores behaved similarly to those from plants without resting spores.

*Phenomena of dehiscence of sporangium and of swarming
and germination of the zoospores*

Discharge. A single apical papilla is formed before the spores are differentiated within the sporangium. This papilla is a thicker and more refractive region of the wall and projects into the sporangium like a plug (Fig. 2*a*). If the contents of the sporangium are plasmolysed, two distinct parts of the plug can be made out; one part remains attached to the wall, while the other is withdrawn with the protoplasm. Barrett⁽²⁾ made a similar observation on the doubleness of the plug in *B. strangulata* (*Allomyces*), and suggested that the inner part of the plug gives rise to the vesicle membrane.

The formation of a vesicle, into which pass the first few zoospores that emerge from the sporangium, was reported by von Minden⁽¹⁴⁾; and I have always observed zoospores to emerge in this manner. The first zoospores to be liberated are seen as a small nearly spherical mass at the apex of the sporangium (Fig. 2*b*); the thin vesicle membrane can barely be distinguished, but attached to it is a small plug, which is probably the outermost part of the original plug or papilla of the sporangium. As more zoospores pass in, the spherical vesicle swells, becoming pyriform (Fig. 2*c, d*). Meanwhile the plug becomes less and less definite until finally it is no longer distinguishable, and the whole plug appears to have been absorbed into the vesicle membrane. When the vesicle is fully expanded, it bursts and the zoospores gradually separate and swim away from the mouth of the sporangium (Fig. 2*e, f*). In one observation (represented diagrammatically in Fig. 2*b-g*), the vesicle reached its full expansion and most of the zoospores separated in three minutes; some of the zoospores, which had entered the vesicle, did not swim away, and no more zoospores emerged from the sporangium.

In general the zoospores which remain in the sporangium have been observed to squeeze out singly by their own movement (Fig. 2*h*). The very active zoospores near the centre of the sporangium push the

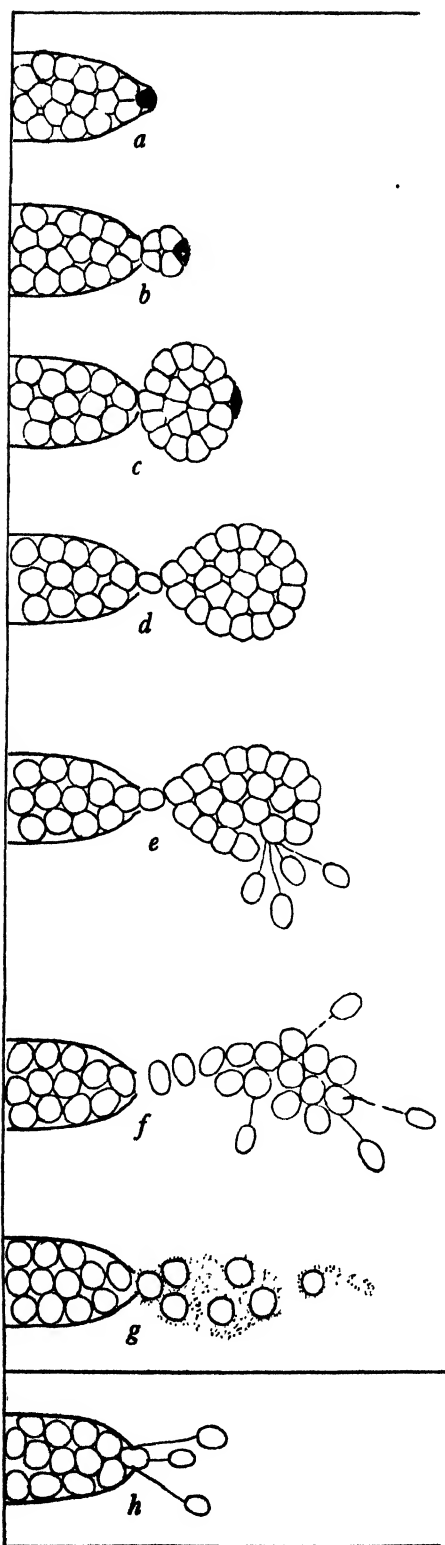


Fig. 2. *a*, the tip of a ripe sporangium. *b-g*, the tip of a sporangium drawn at intervals during the liberation of the zoospores in a vesicle: a process lasting about three minutes. *h*, the tip of a sporangium liberating the zoospores one by one.

less active ones through the opening. The zoospore is egg-shaped and the more pointed end bearing the single flagellum issues last from the sporangium. The flagella of emerging zoospores frequently become entangled and zoospores may be seen tugging and swinging on their flagella, which are about four times as long as the spore.¹ Finally they become free from one another and may swim continuously for about two hours, but the duration of swarming varies considerably.

Swarming. Motile zoospores of *B. Pringsheimii* were first described by Thaxter (22) as oval or elliptical, and generally biflagellated, the two flagella arising side by side from the smaller end of the spore. He described the nucleus as very large and "sub-triangular" in outline, its base connected with that of the flagellum by a fine strand of cytoplasm. Petersen (16) described the zoospores as uniflagellate and von Minden (14) considered uniflagellate zoospores to be typical, while Kanouse (9) described forms with one, two and three flagella. Cotner (4) has shown that they are uniflagellate and uninucleate when allowed to develop normally under optimum conditions, but at temperatures above 14° C. and below 11° C. incomplete cleavage of the protoplasm occurs, so that a number of giant spores are formed with two to three nuclei and a corresponding number of flagella.

In this investigation zoospores have been observed to range in size from 5×6 to $6 \times 9 \mu$. As these observations were made on living material, the number of nuclei could not be determined, and it is possible that the larger zoospores may be the abnormal forms which Cotner has described. The zoospore moves in a fairly uniform manner and an asymmetric spore may be seen to rotate on its longitudinal axis. At any time after emergence a zoospore may show amoeboid movements, while other zoospores are still swarming, but frequently there is no amoeboid stage.

Fields of motile zoospores have been watched to see if zoospores from different sporangia or from sporangia which are borne on different plants show any tendency to fuse. The zoospores vary somewhat in size and some zoospores are less active than others, but from their behaviour there has been no suggestion of a larger female and a smaller male gamete. The flagella of two zoospores have frequently become intertwined; they have, however, always separated later by their own tugging or by the intervention of a third zoospore. Two zoospores have often been seen to come to rest side by side and undergo amoeboid movements, and then one or both have swum away. It is possible that the right combination of gametes (if they are such) has not been obtained, but as many fields of mixed spores have been watched and as germination without fusion has been seen in no less than thirty-two of these it seems unlikely that the zoospores are gametes.

¹ Sparrow (19) has described a similar phenomenon in *Monoblepharis*.

Germination. The early stages of germination have been followed in zoospores which have swarmed and germinated in pond water or soil-extract solution on a slide, while later development has been observed in these solutions only on an agar medium. Similar sporelings and young plants have been found on tomatoes collected from the ponds and from the aquaria. After two to seven days the tomato skin becomes covered with numerous colourless spots, which may be groups of sporelings, or they may be bacterial zooglea. Some days later uninucleate spores have been observed within the zooglea and a few days later again plants of *Blastocladia* have been recognized. Since heavy bacterial zooglea were not always first developed it is unlikely that they play much, if any, part in the foundation of the pustule, yet they do not appear to hinder the development of the sporelings or young plants.

Soon after the zoospore comes to rest a small protuberance, which is the beginning of the germ tube, appears from the spore. The germ tube grows rapidly, while there is little, if any, development of the main body of the spore (Fig. 2*a-d*). Then, when the germ tube is well established, the main body of the spore enlarges and the nucleus divides (Fig. 2*e*). From the subsequent development it is clear that this germ tube is the initial of the rhizoidal system and that the main body of the sporeling enlarges to give the primary main axis. The nuclei in this main axis divide many times while the axis elongates, lobes and begins to branch (Fig. 3*e-p*). Although simultaneous division of the nuclei has been observed in later stages, the nuclei must at first divide independently, as sporelings frequently contain an uneven number of nuclei. Meanwhile the rhizoid elongates and widens, but it appears to branch very little or not at all, until the main axis is further established, when it must branch freely, since it is an elaborate system in mature plants.

SUMMARY

1. *Blastocladia Pringsheimii* is shown to exhibit wide variation in habit, but no *periodicity* in either growth-form or in frequency of resting spores and "hairs".

2. The development of the sporangium with simultaneous divisions of the nuclei is described.

3. The following facts are demonstrated for the zoospore.

(*a*) The swarmers are shown to emerge in a vesicle under certain conditions.

(*b*) No fusion has been observed between the swarmers, and it is concluded that they are asexual and rightly termed zoospores.

(*c*) The zoospore germinates by a small tube, which gives rise to the primary rhizoidal system, while the main body of the spore expands into the main axis.

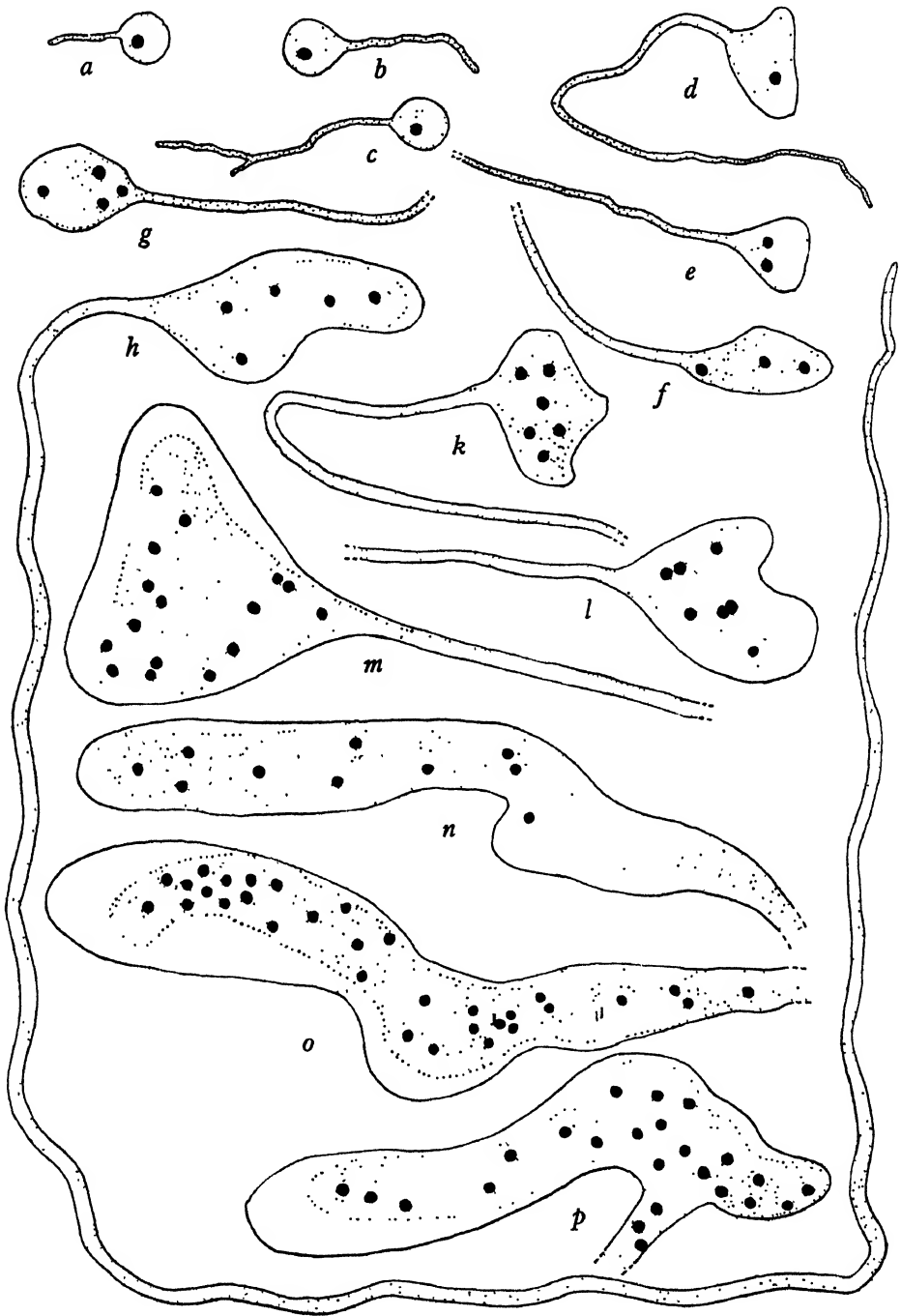


Fig. 3. Germlings ($\times 1000$). In the older ones only the emergent part is shown the rhizoidal part being already extensively developed.

In conclusion, I should like to thank Miss E. M. Blackwell for her helpful criticism and stimulating interest throughout this research and Mrs Topping (Dr M. P. Hall) for suggesting the subject.

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THE LIFE HISTORY AND MORPHOLOGY OF *DICRANOPHORA FULVA* SCHRÖT*

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(With Plates X and XI and 80 Text-figures)

THIS member of the Mucorales was first found by Schröter near Baden in 1877, and again in 1879, growing on *Paxillus involutus*. In 1904 it was found on *Gomphidius viscidus* in the Peloponnese and sent to Vuillemin⁽²⁾, and in 1927 Ling-Young found it on *Boletus scaber* in the forests of the Puy de Dôme. In addition, an unnamed species of *Dicranophora* found by Thaxter on Boleti at Kittery Point, Maine, U.S.A., was briefly described by Blakeslee in his classic paper of 1904. These are the only recorded appearances of the fungus in nature.

The accounts are short and incomplete and differ on a number of points. Blakeslee and Ling-Young were concerned only with the sexual stage. Vuillemin, whose paper is the only one published on *Dicranophora* alone, dealt chiefly with the sporangial stage. The chief characteristics of the fungus, as described by Schröter⁽⁵⁾, may be summarized thus: Protoplasm yellowish red. Sporangia of two types, (1) large, with conical columella, and small elliptical spores, and (2) small, with two- or three-pointed forked columella and one or two much larger reniform spores. Zygosporangium spherical, chestnut-brown, smooth or with fine warts, suspensors very unequal, the one swollen, the other thread-like.

Reference to Vuillemin's paper⁽²⁾ shows that he regarded the supposed columella as "rudimentary branches of the apophysis, continuing the dichotomy of the axis, and having nothing to do with a columella". The saddle-shaped "apophysis" cannot be called a columella, as it does not encroach upon the interior of the sporangiole. He compares it, however, to the subsporangial swellings of *Pilobolus* and the Entomophthorales, since, he states "le redressement brusque du plancher en forme de selle a pour effet de décoller le microcyste et de le projeter avec force". Flies, he suggests, probably assist dissemination.

Schröter⁽¹⁾, in describing the copulating branches, states that the upper third of the thicker one is distinguished by a cross-wall for the formation of the zygosporangium, and remarks that their unlike character "points distinctly to a sexual difference in the branches, and suggests an antheridium and an oogonium". Blakeslee's figures⁽⁴⁾, Pl. I,

* Part of thesis approved for the Degree of Doctor of Philosophy in the University of London.

figs. 10–14) show the development of a “bulge” on the female hypha, in contact with the male, which gives rise to the entire gametangium, and is much more clearly demarcated from the parent branch than that shown by Ling-Young (3), who was, however, in agreement with him that “L’ampoule copulatrice du sexe femelle ne se forme que sous l’action provocatrice du gametophore mâle”. Gwynne-Vaughan & Barnes (7) reprint in their textbook two of Blakeslee’s figures, and state that one of the gametangia “becomes swollen, and a characteristic bulge develops on its stalk”—a point not mentioned by Blakeslee, and which seems to require further explanation. The zygospore coat is also variously described as smooth or finely warted, by Schröter, and characterized by deep crevices and resembling that of *Spinellus*, by Vuillemin.

The systematic position allotted to *Dicranophora* by Schröter (5) and adhered to in later systematic works, is next to *Thamnidium*, from which it is distinguished by the presence of a columella, and of a peculiar type of spore, in the sporangiole. Blakeslee, and Gwynne-Vaughan and Barnes, referring to the sexual stage, state that “the condition is essentially similar to that in *Zygorhynchus*”, and Vuillemin, despite his denial of one of the chief differences from *Thamnidium*, considers that it is not related to that genus, but should be placed between *Sporodinia* and *Spinellus*. Finally Fitzpatrick (6), although taking into account Vuillemin’s work, nevertheless places *Dicranophora* in the Thamnidiaceae, and distinguishes it from *Thamnidium* only by the claw-like branches clasping the sporangiole, and the dimorphism of the spores in the two types of sporangium, which, he states, makes the genus of unusual interest.

It is clear, therefore, that a thorough investigation of the life history and morphology of this fungus is needed.

MATERIAL AND CULTURAL METHODS

The strain of *Dicranophora fulva* Schröt. which I have studied was originally collected by Ling-Young and sent in 1928 to the Centraal-bureau voor Schimmelcultures, Baarn, Holland, whence it was obtained in 1934. During the two years that I have successfully kept it in culture it appears not to have changed its character, although it is extremely variable, and sensitive to conditions of temperature, light and humidity.

Sensitiveness of the fungus

(1) *To temperature.*

A culture placed in an incubator at 26° C. for twenty-four hours was killed outright. Another, subjected to a day temperature of about 23° C. for a week-end (sixty hours), gave only one viable sub-culture out of ten attempts, and this was a sterile variant with spore-

less sporangia. The optimum temperature for growth and sporing is about 19° C. Exposure to freezing temperatures was found to have no injurious effect.

(2) *To light.*

Cultures left on the laboratory bench developed sporangia, but no zygospores. So also did those kept in the dark and examined daily. The extraordinary sensitiveness of the fungus to light was not discovered until plates left in the dark without examination for a week were found to bear a mass of zygospores, but no sporangia. It was then found that sporangia are formed only where there has been some illumination, whereas the early stages of zygospore formation occur only in the dark. A very slight exposure to light, e.g. that caused by a brief examination of the culture, is sufficient to induce sporangial formation, and to inhibit the development of the larger zygothoric branches, which lose their contents into the surrounding mycelium, and are left as empty sacs.

Although this sensitiveness to light has not been referred to by previous authors, and it is possible that it is a pathological condition associated with long culturing, there is reason to suppose that Blakeslee's material may have shown the same thing. He states that, of three stender dish cultures, "one left covered in a drawer produced abundant zygospores and few sporangia; while the other two, one of which was placed uncovered on a laboratory table, and the other in a sealed vessel with calcium chloride, ... produced sporangiophores but no zygospores". He attributes this to the difference in humidity, but the result is that which would be obtained by light sensitivity, assuming that the "sealed vessel" was of glass and was kept in the light, as would be probable. The few sporangia on the covered plate could be induced by once examining it. Blakeslee also states that "under favourable conditions" sporangia and zygospores are produced on the same plate. This also is compatible with light sensitivity, since this result is obtained when, as is customary, the plates are kept in a dark incubator which is opened at intervals. Zygospores and sporangia are also obtained together on a culture grown in very diffuse daylight, which accounts for the occurrence of zygospores on the underside of the cap of an Agaric, and between the interstices of a piece of bread. Probably the night is a sufficiently long period of darkness to allow the completion of the earliest and most sensitive stages. A more detailed investigation of this phenomenon is now in progress.

(3) *To humidity.*

Differences in humidity do not appear greatly to affect the relative numbers of sporangia or zygospores. The fungus, however, is killed

by desiccation, and the older white parts of the aerial mycelium are not viable. Hence subculturing every ten days is necessary for safety. In a saturated atmosphere the sporangiophores tend to produce irregular swollen masses (e.g. Text-fig. 10), the conditions for the formation of which require further investigation. When the culture is flooded the sporangiophore initials burst, and extrude all their protoplasm as a long coil. In several cell cultures, however, in which a thin film of condensed water was formed, the swollen tips produced a number of radially arranged "buds", the whole resembling, in general arrangement, the oedocephaloid "head" of *Cunninghamella* (Pl. X, fig. 24), although differing in many details, and particularly in the fact that the "buds" do not drop off, but may germinate *in situ* as shown in the photograph. Blakeslee's *Mucor* I which "produces chlamydospores which bud in a characteristic oedocephalum-like manner" (4), probably affords a closer comparison, although in *Dicranophora* chlamydospores are never formed.

Media

In order to discover what type of medium was most suitable, I grew the fungus on a range of common media, under comparable conditions of light and temperature. The figures given in the following table are purely relative, and were assigned on an estimate of the appearance of the cultures under a low power binocular. Greater accuracy is impossible without rigid standardization of all conditions.

Medium	Mycelium	Yellow pigment	Sporangia	Zygospores
Plain agar	1	0	0	0
Glucose asparagin agar	3	0	1	0
Glucose peptone agar	5	0	2	0
Potato agar	9	4	10	2
Prune agar	10	10	10	3
Malt agar, pH 6	6	6	8	7
Malt agar, pH 7.5	7	7	7	8
Malt broth, pH 7.5	8	7	2	10
Brown bread	.	8	9	6

It will be seen that no zygospores, or pigment, are formed on the artificial media, and that malt agar is the best general purpose medium. 3% malt agar, cleared and filtered for the sexual stages, was accordingly used for the bulk of the work. Difference in pH has no great effect, but on the whole alkalinity appears to favour zygospore production. A fairly vigorous growth is, however, obtainable on this medium from pH 4 to 8. Malt broth does not produce sporangia until a thick mat of mycelium has made a skin capable of bearing aerial branches.

An attempt to culture *Dicranophora* on its natural host, *Boletus scaber*, failed owing to bacterial infection, but its luxuriant growth on brown bread, with superficial sporangia, and zygospores between the folds of

the crumbs, probably resembles the natural state (see Ling-Young (3)) much more closely than do the plate cultures.

Technique of culturing

For the study of the sporangial stages Vernon's technique (8) was at first adopted, but the rapid drying which takes place on the microscope stage was a serious difficulty, and it was found more satisfactory to cut out blocks of agar, bearing mycelium from a plate culture, and put them under a cover-slip on an ordinary ring cell. Sporangiphores then begin to emerge from the agar on the second day, and grow parallel to the cover-slip, producing sporangia usually on the third day when the room temperature is fairly high (about 17° C.). The chief trouble in such a closed cell is excessive condensation, which is very liable to spoil serial observations, as may be seen from the series of photographs (Pl. X, figs. 14-23) of which the later stages are partly obliterated by a creeping film of moisture. To overcome this difficulty a type of split cell was improvised which is described in the section on the sporangial stage.

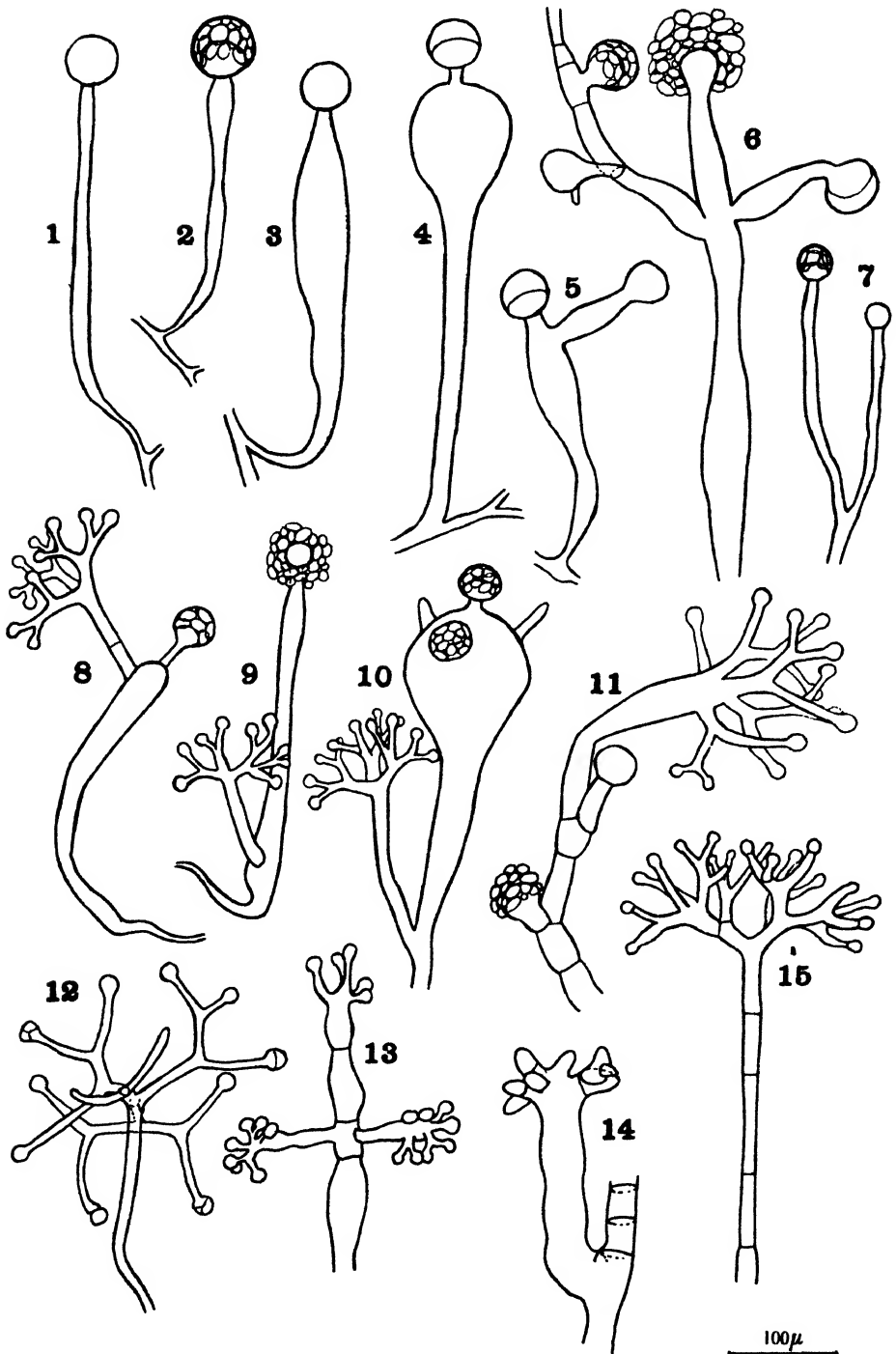
The sensitiveness of the sexual stage to light necessitated a special technique for the study of the living material. A range of light filters from ultra-violet to red was interposed on the substage of the microscope, but the sex organs always lost their contents and ceased development after the exposure necessary for a camera lucida drawing. Finally I found it possible, by using a deep red dark-room filter on the lamp, and a pale green filter on the substage, to make hurried outline sketches of the sex organs in the dim orange light resulting, without always interfering with their development. When it was necessary to focus on details the green filter was pushed aside and replaced as soon as possible. In this way the complete development from early contact to zygosporangium was followed on five occasions only, out of several hundred attempts. Even less success rewarded attempts to take serial photographs. Pl. XI, figs. 36, 37, are the only two successive photographs showing further development of the sexual stage. The exposures were fifteen seconds on a hypersensitive panchromatic plate, with a 60 W. bulb.

It was found necessary to design a special type of chamber which would allow examination of the sexual stage with the 1/6 in. objective without the necessity of disturbing the culture. Details of this have been published elsewhere (9).

THE SPORANGIAL STAGE

(1) *The sporangiophore*

A culture of *Dicranophora* grown in the light exhibits a variety of sporangial forms, some of which are shown in Text-figs. 1-15. The variation in appearance is chiefly due to the sporangiophore, which



Text-figs. 1-15. *Dicranophora fulva*. Forms of the sporangiophore.

may be simple or branched, narrow or swollen. The commonest type of simple sporangiophore has the slightly swollen form shown in Text-fig. 2, but the swelling may be absent (Text-fig. 1) or much more pronounced (Text-figs. 3, 4). The large subsporangial swelling shown in Text-fig. 4 has a superficial resemblance to that of *Pilobolus*, which is increased when the short stalk between it and the sporangium is absent, as sometimes occurs. It has, however, no explosive function, and has even been pricked with a glass splinter without collapsing. Text-fig. 10 shows one such swelling which bears several sporangia, and is also putting out two branches.

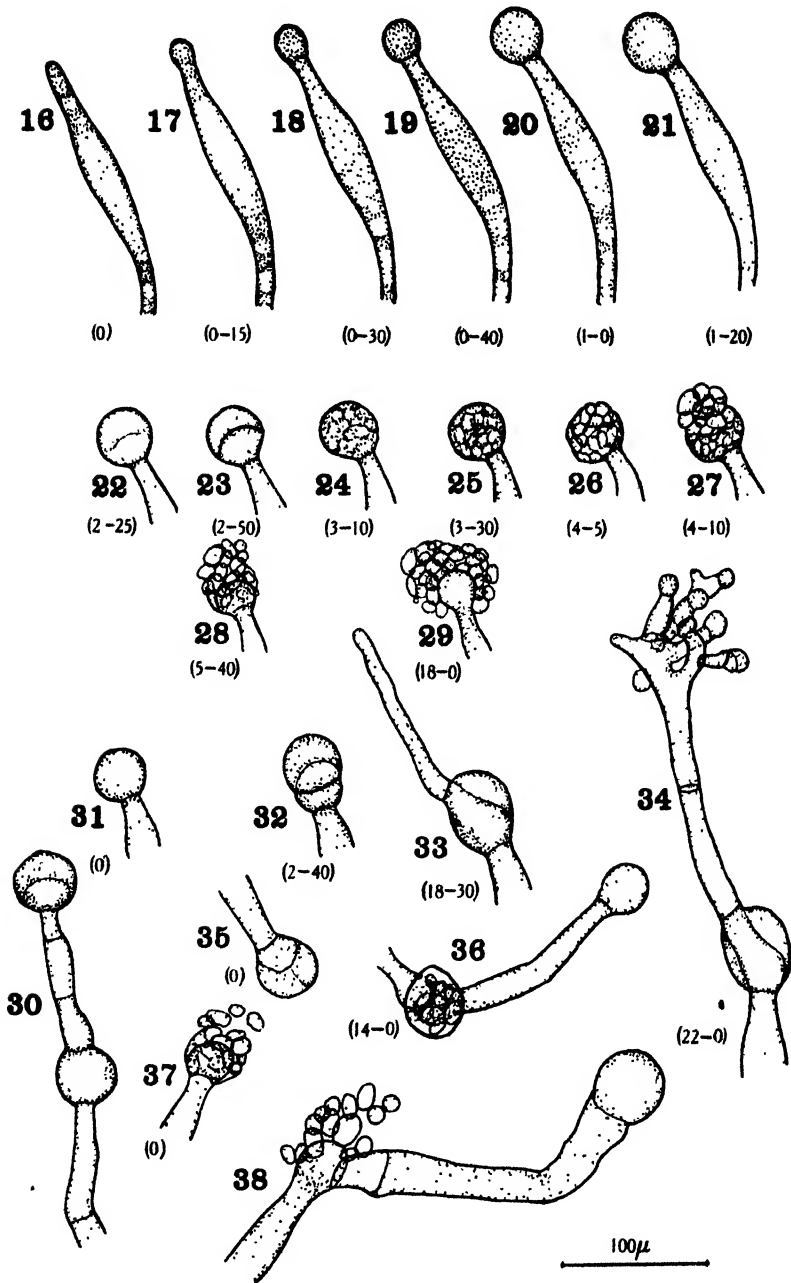
The large sporangia are most frequently borne on unbranched sporangiophores, but Text-figs. 5–7 show that slight irregular branching may occur. They may also be associated with the sporangiole branch systems (Text-figs. 8–11), but more commonly not. The main axis of the sporangiole branch system may, like the sporangiophore of the large sporangium, be thin (Text-fig. 15) or more or less swollen (Text-fig. 14); and the branching may be lax (Text-fig. 12) or dense (Text-fig. 13), varying chiefly with the light intensity. Forms intermediate in size and in appearance between the typical sporangiophore and sporangiole branch system, are sometimes to be found (e.g. Text-fig. 7).

(2) *The sporangium*

Text-figs. 16–29 illustrate, by a series of camera-lucida drawings, the development of a single sporangium. In the early stages dense granular protoplasm is visibly flowing in at a rapid rate, and pressing up the club-shaped sporangiophore in the form of dense “plugs” which tend to coalesce towards the tip. A wide area of mycelium is thus drained of protoplasm in a short time, the highest rate of flow measured being 115μ in ten seconds. The tip swells to the full size of the spherical sporangium in about an hour, and a back flow of protoplasm down the sporangiophore has been seen after the maximum size has been attained.

The dense protoplasm which packs the swelling tip usually ends in an irregular line visible just below the line of demarcation between the curved surface of the sporangium and the cylindrical sporangiophore. A septum has not been seen in this position, nor in the position figured by Schröter ((5), Fig. 113B) a little farther down the stalk, although septa are sometimes found in the branched sporangiophores.

A delay of about one hour occurs between the attainment of full size and the first appearance of the columella as a faint division across the sporangium (Text-fig. 22). For a short period the columella boundary is visible as a clear zone often appearing nearer the top of the sporangium, and apparently, owing to the effects of refraction by the curved surface, extending to the sides rather than the base



Text-figs. 16-29. *Dicranophora fulva*. Stages in the development of a single sporangium.

Text-fig. 30. Branch produced from a young sporangial swelling.

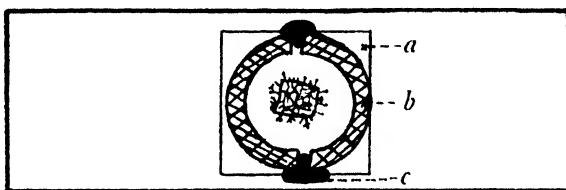
Text-figs. 31-4, 35-6, 37-8. Three different series showing sprouting of the columella.

The numbers in brackets show the times, in hours and minutes, from the previous figure marked (0).

(Pl. X, fig. 4); this boundary, however, becomes indistinct at the onset of spore formation and remains so until the spores are fully formed. The particular sporangium shown in Text-figs. 16–29 burst in contact with the cover-slip after about four hours, the spores flowing out to form a patch, for the most part in the plane of the cover-slip, around the columella (Text-fig. 29 and Pl. X, fig. 8).

When the sporangium does not touch the glass, it gives place to a large water drop which contains the spores (Text-fig. 2 and Pl. X, fig. 2). This may be compared with the oidial drops of *Coprinus* described by Brodie⁽¹⁰⁾. The thin smooth hyaline membrane is invisible, except in mounted material, and it is impossible by direct observation to determine exactly when the sporangial wall bursts and is replaced by the water surface, but if the drop is allowed to dry, it is possible to see whether it keeps its shape or resolves itself into a mass of spores.

For this purpose a type of cell (shown in plan in Text-fig. 39) was improvised, in which it is possible to control the rate of drying of the



Text-fig. 39. Plan diagram of a type of split ring cell used for observations on sporangia. *a*, square cover-slip; *b*, split glass ring; *c*, wax plug.

contents to some extent. A small block of agar cut from a plate culture is placed under a cover-slip which is vaselined to the split glass ring. The gaps in the ring are at first blocked with wax, but when the sporangial drops have appeared the humidity is lowered by removing some of the wax and allowing the water vapour to escape. By watching a number of marked sporangia it was possible to determine that no membrane was present in most of the "sporangial drops", since they dried down into irregular spore masses. In three, however, out of twenty observed, the membrane was still present. One of these was seen to contract from 50 to 38 μ in diameter in two hours—a loss of about 50 % of its volume, and 42 % of the surface area of the membrane, which must therefore be extremely elastic. On two cultures on an old malt agar, the sporangial membrane, which elsewhere was invariably quite smooth and hyaline, was covered with crystals, probably of calcium oxalate, since they did not dissolve in dilute mineral acid (see Pl. X, fig. 11).

A sporangium, or at least a terminal swelling entirely similar to a young sporangium, may give rise to a hypha without further develop-

ment (Text-fig. 30). On the other hand, the columella may sprout before or after spore formation. The sporangium shown in the series Text-figs. 31-34 gave rise in two and a half hours to the remarkable two-celled form shown in Text-fig. 32, in which, apparently, the sporangial cavity encloses only the top half of the columella. The lower cell gave rise overnight to a branch which emerged near the top of the upper cell (Text-fig. 33) and produced a bunch of sporangioles in a few hours. Such two-celled forms have been observed a number of times, and may sprout from either cell, though more commonly the basal one. The distal cell has only once been seen to form spores.

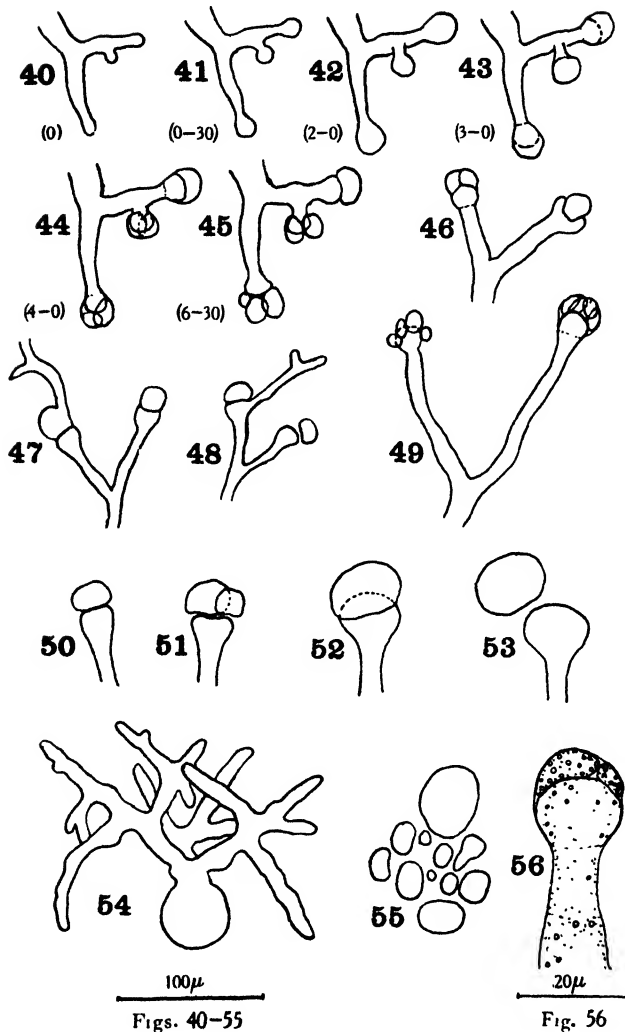
Sprouting of the columella after spore formation is shown in Text-figs. 35, 36, where the branch has pushed aside the sporangial sac in which spores have nevertheless been formed, and in Text-figs. 37, 38, in which the sporangium has burst before the appearance of the branch. Pl. X, fig. 13 is a photograph of the same object as Text-fig. 38, taken a little later. The columella now appears merely as a "knee" in the sporangiophore, but can be distinguished by the patch of spores beside it.

From the figures (particularly Pl. X, fig. 2) it will be seen that the wall of the columella is more or less spherical. The range of variation is very small—from a slightly depressed to a slightly raised oval.

(3) *The sporangiole*

The development of the sporangiole shows the same stages, and takes about the same time, as that of the sporangium, and it is sufficiently illustrated by Text-figs. 40-45, Pl. X, figs. 3-5 and figs. 14-23. The young sporangiole is more or less globose, although later the membrane adapts its shape to that of the spores beneath, so that it is often two- or three-lobed (Text-fig. 46). The columella is not easily seen owing to the thinness of the scarcely visible membrane which is pressed tightly to it except where separated from it by the spores. Text-fig. 56 and Pl. X, fig. 9, however, place this point beyond doubt. The columella can be clearly demonstrated when sporangioles are mounted in water and gently crushed under the cover-slip. In a small proportion of those which remain unburst, the sporangial membrane separates from the columella and spores (as shown in Pl. X, fig. 9).

The columella is of the same shape as in the large sporangium, namely, a slightly depressed oval, although in mounted material it is frequently distorted. It has the same ability to branch, and "two-celled" forms have also occasionally been seen (Text-fig. 47). On the other hand, the object shown on the right of Text-fig. 46 is the only approach seen to the "tong-like claws" reported in this genus.



Text-figs. 40-5. *Dicranophora fulva*. Stages in the development of a small part of a sporangiole branch system.

Text-fig. 46. "Lobed" sporangiole and (right) claw-like outgrowths of columella.

Text-fig. 47. "Two-celled" sporangioles, showing sprouting of distal cell (left).

Text-fig. 48. Sprouting columella of sporangiole.

Text-fig. 49. Small "intermediate" sporangia.

Text-figs. 50-1. Large sporangiole-like forms (after disappearance of the membranc).

Text-figs. 52-4. The liberation and germination of a "giant" spore.

Text-fig. 55. Group of spores showing variation in size.

Text-fig. 56. Sporangiole in section, showing columella, two spores and membrane.

Camera lucida drawing under 1/12 in. objective.

The numbers in brackets show the times, in hours and minutes, from the previous figure marked (0).

This was found in a culture grown on bread. The columella is concave, and clasps the single spore, but the effect is probably exaggerated by mounting in water.

(4) *The spores*

In point of size and spore number, some of the largest sporangioles are indistinguishable from small sporangia (e.g. Text-fig. 49). A count of the spore numbers in five hundred burst sporangioles in mounted material gave the following frequencies:

Spore numbers	1	2	3	4	5	6	7	Over 7
Frequencies	264	156	64	13	1	1	1	0

For this purpose any spore-bearing body borne on a dichotomous branch system was reckoned a "sporangiole". On the other hand, three out of a hundred "sporangia" borne on simple sporangio-phores were found to have less than ten spores (e.g. Pl. X, fig. 7), and forms borne on very lax branch systems (Text-fig. 7) were quite impossible to distinguish as one or the other. Very large "sporangioles" with one or two spores only (Text-figs. 50, 51) provide further links, and the discovery of large one-spored sporangia completes the evidence as to the continuity of the two forms.

The range of spore size in sporangia is extremely wide, normally from 5 to 30 μ , but I was surprised to find in some spore mounts "giant" spores 40–50 μ in diameter, i.e. the full size of an average sporangium. The problem was solved by chance manipulation of a "two-celled form" in which the distal "cell" was detached with a glass splinter (Text-figs. 52, 53) and transferred to a malt-agar slide, where it germinated as shown in Text-fig. 54. Some difficulty was found in pulling the giant spore away, for it sprang back to the columella several times under the pull of the elastic membrane, which was invisible under the low power used. A similar giant spore (not quite so large—about 30 μ in diameter) is shown standing on a flattened columella in Pl. X, fig. 6. This, however, despite its size, was borne on a lax dichotomous branch system.

It is evident, therefore, from the foregoing observations, that the distinction between sporangium and sporangiole is quite an arbitrary one, and the typical sporangiole does not differ from the sporangium, except in size, spore number, and position on a branchlet farther from the protoplasmic stream of the main axis, which would account for the other differences.

The alleged dimorphism of the spores in the two sorts of sporangia is sufficiently disproved by the average dimensions given below, which are based upon measurements of two hundred spores from each sort,

arranged in frequency classes of $1\ \mu$. The corresponding figures for *Mucor hiemalis* are given for comparison.

	Length	Breadth	Ratio length to breadth
<i>Dicranophora</i> , sporangium spores	Mean $13.46\ \mu \pm 0.82$ Range $5-30\ \mu$	Mean $10.83\ \mu \pm 0.77$ Range $3-25\ \mu$	1.24
<i>Dicranophora</i> , sporangiole spores	Mean $12.95\ \mu \pm 0.68$ Range $5-20\ \mu$	Mean $9.23\ \mu \pm 0.65$ Range $3-20\ \mu$	1.40
<i>Mucor hiemalis</i>	Mean $5.40\ \mu \pm 0.42$ Range $3-8\ \mu$	Mean $3.40\ \mu \pm 0.34$ Range $2-6\ \mu$	1.59

It is clear from these figures that the spores in the sporangioles are not larger than those in the sporangia, and, in fact, appear to be slightly smaller, although the differences between the mean dimensions are not significant.

The comparison with *Mucor hiemalis* serves to emphasize the extraordinarily wide range in spore size of *Dicranophora* (see Text-fig. 55 and Pl. X, fig. 7). Spores at the top of the normal range ($25 \times 30\ \mu$) are as much as 417 times the volume of the smallest spores ($3 \times 5\ \mu$). In a single sporangium chosen at random the largest spore ($14 \times 18\ \mu$) was found to be seventy-eight times the volume of the smallest ($3 \times 5\ \mu$). In contrast, the largest spore of *Mucor hiemalis* was found to be less than twenty-two times the volume of the smallest, and in a single sporangium, about seven times. It seems probable that, in regard to spore size, *Dicranophora* is by far the most variable of the Mucorales.

The absence of the largest spores from the sporangioles, as indicated by a smaller range, is responsible for the lower mean of the sporangiole spores, and the fact that only typical sporangia and sporangioles were chosen for these measurements would tend to this result, by excluding forms which might be regarded as large sporangioles or small sporangia, and might be expected to contain spores of corresponding size.

The slightly greater ratio of length to breadth in sporangiole spores, corresponds with a higher proportion of long narrow spores in the sporangioles, in which the spores are necessarily compressed between the membrane and the columella (Text-fig. 56), and consequently tend to have a convex outer, and a concave inner surface, thus assuming a kidney shape which is most obvious where there is only one spore in the sporangiole. When liberated, however, they round off to an elliptical shape. Similar, but less obvious, compressed spores are found among the sporangiospores, and it is therefore clear that the spores in the sporangioles differ from those in the sporangia only in the slightly more compressed shape imposed upon them by the limited curved boundaries of the sporangiole.

(5). *The "sporangial unit"*

The fact that the young sporangiophore, before it swells at the tip, is indistinguishable from the sporangiole branch system initial, before it has started to dichotomize, suggests that the organs produced by them are homologous. An attempt was therefore made to compare the numbers of spores produced by the large sporangium and the sporangiole branch system, respectively.

The number of spores in a sporangium may be fairly easily counted when the sporangium bursts against the cover-slip, forming a patch of spores about one deep, held together by the surface tension of the water (Pl. X, fig. 8). It is possible to exclude spore patches which are incomplete, or have received spores from other sporangia, and for the purpose of the count only sporangia on an unbranched stalk were included. In this way a count was made of the spore numbers in a hundred sporangia, chosen at random from twelve different cultures in ring cells.

It was impossible, however, to count directly the number of spores in a sporangiole branch system. The sporangioles therefore were counted, on a hundred branch systems on the same twelve cultures, and their number multiplied by a mean obtained by counting the number of spores on two hundred burst sporangioles in the same material.

The results obtained from these counts were as follows:

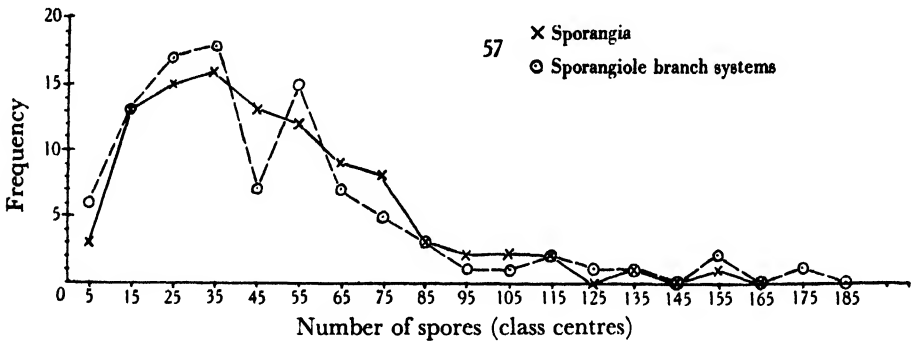
Spore numbers in	Range	Mean	S.E.
Sporangia	5-160	47.5	6.82
Sporangiole branch systems	4-180	46.8	? Large

The closeness of the means is of little significance in view of the large unknown error introduced with the additional estimate involved in the second mean. On the other hand, the great similarity of range and form of the two frequency curves (Text-fig. 57) cannot be put down to chance, and it seems evident that the initial club-shaped branch gives rise to a "sporangial unit" which, whether it forms one sporangium, or many sporangioles, contains the same amount of protoplasm, and produces about the same number of spores. The factors which determine whether the tip will swell, or branch, remain to be discovered, but there are indications that light is the most important.

(6) *Comparison with previous accounts*

The foregoing description differs most obviously from that of Schröter(5), both regarding spore size, and the conclusion that the two types of sporangium are not essentially different. Schröter describes the sporangiospores as: "etwa 7-14 μ lang, 4-10 μ breit

(meist $7\mu \times 5\mu$)” and the spores in sporangioles as “gross, nierenformig, 22–26 μ lang, 11–15 μ breit”. Not only are these figures in disagreement with mine as regards dimorphism, but it is clear, also, that the variation in spore size, which was remarked upon by Schröter, is even more pronounced than he thought. The difference in shape of the columella, which in my material varies very little from its spherical shape, and never approaches the conical, is probably of less significance, in view of Vuillemin’s statement that it may also be pyriform or hemispherical. It would seem, however, that the process of culturing to which my material has been subjected has selected a strain which is very constant for this factor.



Text-fig. 57. *Dicranophora fulva*. Graph showing the frequency of spore-numbers in one hundred sporangia, and in one hundred sporangiole branch systems, the former being counted directly, the latter arrived at by counting the number of sporangioles per branch system, and multiplying by a mean spore-number obtained from a count of two hundred sporangioles in the same material.

The confirmation of the presence of a columella in the sporangiole necessarily conflicts with Vuillemin’s view, but his interpretation of the “claws” as branches continuing the dichotomy of the axis is supported to some extent by my observations on the branching of the columella, although it cannot be fully confirmed, owing to the absence of these claw-like outgrowths (except for one isolated record) from my material. In view of the fact that Vuillemin saw no living specimens, and thought that the sporangioles themselves could be shed, or even shot off, it seems probable that he missed seeing the very inconspicuous sporangial membrane, and, sometimes at least, mistook the spore for the sporangiole.

THE SEXUAL STAGE

(1) *Early stages in conjugation*

The difficulty of studying the earliest stages of conjugation in *Dicranophora* has been remarked upon by Blakeslee(4). An attempt to reconstruct the process from fixed material led to quite erroneous conclusions, and despite the light sensitivity it was found necessary to make a developmental study of the living sex organs.

My observations are best described by reference to the drawings in Text-figs. 58–80, and the photographs on Pl. XI. The female sex organs arise as swollen sacs, 60μ or more broad, which may be single (Pl. XI, fig. 27) or in groups (Pl. XI, fig. 25) and are densely packed with yellow protoplasm. These may form zygospores at once, or grow indefinitely to form a long, lobed hypha, constricted at intervals, and often branching dichotomously (Pl. XI, fig. 26). The method of growth is peculiar, and is illustrated in Text-figs. 58–61. The end of the sac becomes smoothly rounded, with a clearer border, and then puts out a protuberance, or sometimes two, which swells to about the diameter of the parent sac before behaving similarly. Zygospores are formed on the branch wherever male hyphae come in contact with it, but if unfertilized the swollen lobes may give rise to hyphae of normal dimensions recognizable by the wavy course which they take through the medium (Text-fig. 61).

Most of the female sacs form zygospores in the first stage, before any further growth occurs, or else lose their contents to neighbouring lobes on which zygospores are being formed (Text-figs. 74, 75). The series Text-figs. 62–5 shows the development of a group of three sac-like branches (Text-fig. 62*a, b, c*) which, however, did not reach maturity owing to the illumination necessary for the drawings. The lobe *b* was already producing a bulge before contact with a male hypha caused it to swell out laterally to form the female gametangium.

Legends to Text-figs. 58–80.

Text-figs. 58–80. *Dicranophora fulva*. Stages in zygospore formation.

Text-figs. 58–61. Series showing dichotomous growth of a zygophoric branch.

Text-figs. 62–5. Series showing fertilization of three female sacs at different stages of growth.

Text-figs. 66–7. Formation of a zygospore without any outgrowth of the female organ towards the male.

Text-figs. 68–9. Formation of a zygospore on a large curved branch, resembling the type of reproduction in *Zygorhynchus*. The zygospore shown in Text-fig. 69 (as also in Text-fig. 65 and one of those in Text-fig. 67) has not reached maturity owing to the effects of light, and the large gametangium has become empty, except for a mass of protoplasm which blocks the pore in the large septum.

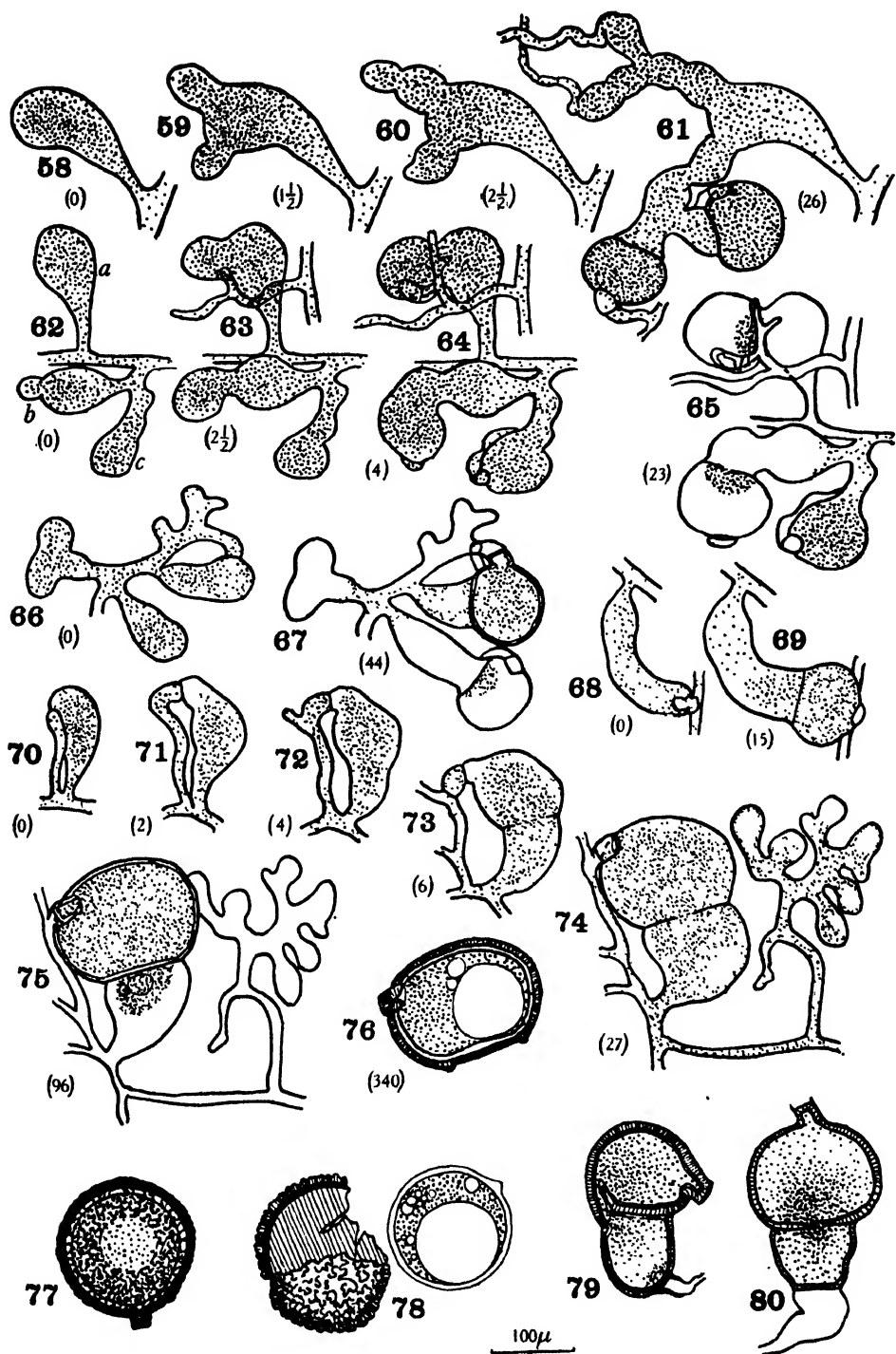
Text-figs. 70–6. Developments of a zygospore with formation of a papilla on the female sac, followed by swelling of the whole organ. The “clear stage” of the male gametangium, which precedes the thickening of the wall, is shown in Text-fig. 74, and the emptying of neighbouring unfertilized lobes in Text-figs. 74 and 75. Text-fig. 75 also shows a mass of oil and granular matter left behind in the large suspensor, and the mature zygospore, a fortnight old, with double wall, large oil drop, and “antheridial boss” is shown in side view in Text-fig. 76.

Text-fig. 77. Mature zygospore, seen from the direction of the large suspensor, the position of which is indicated by the unmarked patch on the creviced wall.

Text-fig. 78. Mature zygospore, crushed to separate the creviced exospore from the thick hyaline endospore. The slight projection on the inner coat fits into the “boss” on the outer one.

Text-figs. 79, 80. Two abnormal zygospores in which the septum cutting off the large gametangium is incomplete. In that shown in Text-fig. 79 the wall of the whole female branch is thickened. In the other (Text-fig. 80) the thickening is limited by a second septum across the branch.

The numbers in brackets show the time in hours from the previous figure marked (o).



Text-figs. 58-80 (for legends see previous page).

A similar bulge on lobe *a*, however, appears to have itself formed the gametangium, while the lobe *c* continued, after contact with the male, to swell at the tip, and would presumably, if left in the dark, have formed a zygosporangium without any outgrowth towards the male, as in fact was observed elsewhere (Text-figs. 66, 67).

Before contact, the male hypha is not differentiated from the other hyphae of the mycelium, and is impossible to distinguish as such; moreover, the light sensitivity prevents direct observation of the growth of the hyphae, but it is evident from Text-figs. 58–65 that the male and female branches are not always in contact from their inception, and it is difficult to interpret figs. 28 and 36 on Pl. XI as anything but early stages in contact.

The rate of linear growth of the male branch appears to be much greater than that of the female, which is to be expected from its smaller diameter. The frequent sharp bending of the male towards the female branch (Pl. XI, figs. 28, 32) suggests some chemotropic influence, which would naturally show its effect more on the faster growing hypha. Sometimes, however, the larger branch in its growth comes against the side of a narrow hypha which puts out a male progametangium at the point of contact (Text-figs. 68, 69). Here the chemotropic effect, if any, must be shown by the female. The figures show (e.g. Pl. XI, figs. 32, 35) that the male branch frequently arises near the base of the female, but may also come from an entirely different hypha.

The type of development shown in Text-figs. 70–6 appears to be the commonest, and resembles more closely that figured and described by Ling-Young. Here the papilla on the large sac is formed in response to contact with the small hypha, but the whole female branch then swells enormously, and the gametangium does not arise from the papilla alone. Pl. XI, figs. 29, 30, show early stages in contact, with a papilla on the female. Pl. XI, fig. 36, shows a very early stage in contact, and Pl. XI, fig. 37, shows the same object the next morning. Although partly obscured by shadows of hyphae and drops which have appeared during the interval, it shows clearly the swelling up of the tip of the male hypha, the passage of granular protoplasm into the tip, and its close application to the female sac, which has swelled out considerably towards the male.

(2) *The passage of contents*

The septum cutting off the male gametangium appears before that on the female. The larger septum of the latter grows in slowly towards the centre, and it is impossible to know by direct observation when it is complete, but the light sensitivity of the fungus can be used to obtain evidence on this point.

When developing zygosporos are illuminated, the granular contents flow out of those gametangia which have incomplete walls, and the extent of the pore can be observed in the empty sacs. Where the pore in the large septum is small it is usually blocked by a mass of protoplasm which remains behind (Text-figs. 65, 67). A number of male gametangia in which the basal septum was complete were seen to empty themselves through the female, and actual movement of granules from one to the other was twice followed. The pore, which is at first an irregular oblong slit, can sometimes be seen when the organs are empty. Evidence of this passage of contents from the male to the emptying female gametangium is provided by the series of photographs (Pl. XI, figs. 38-40).

That passage of contents also occurs under normal conditions can be deduced from the appearance of a clear stage in the development of the male gametangium (Text-fig. 74 and Pl. XI, fig. 32), and is placed beyond doubt by the complete obliteration of its lumen by the developing wall of the zygosporos (Pl. XI, fig. 42).

(3) *The zygosporos*

During the period of swelling a rapid flow of protoplasm into the growing spore is visible along the neighbouring hyphae. This reverses its direction when the septum cutting off the gametangium is complete, and the large suspensor is emptied back into the mycelium, although a mass of granular matter and oil, which eventually disappears, is frequently left behind (Text-fig. 75). In this way practically the whole content of a culture kept in the dark is incorporated in the zygosporos, which tend to vary in size according to the density of the mycelium. Thus, measurements of a hundred in side view (i.e. as in Text-fig. 76) from two different media, gave the following figures:

Medium	Mycelium	Zygosporos per unit field (average of 10 counts)	Mean long axis	Mean short axis	Ratio length/ breadth
3 % malt agar	Dense	44.1	$162.7 \mu \pm 6.0$	$133.9 \mu \pm 5.7$	1.215
Weak cherry agar	Less dense	47.3	$141.9 \mu \pm 5.1$	$116.8 \mu \pm 5.1$	1.215

The size differences are significant. The volume of a zygosporos of average dimensions on malt agar is about 0.0018 c.mm.—sufficient to absorb the contents of about 23 mm. of hyphae.

It is clear from these measurements, and from Text-fig. 76 and Pl. XI, fig. 42, that the zygosporos is not spherical, but is somewhat flattened on the side of the large suspensor, and is, in fact, rather the shape of a horse-chestnut seed. The outer coat is deeply and irregularly creviced (Pl. XI, fig. 43) except for a patch in the position of the

large septum (Text-fig. 77). The thick hyaline inner coat is easily separated from the outer by crushing (Pl. XI, fig. 44) and bears a small beak (Text-fig. 78) fitting into the boss on the outer wall which is all that remains of the male gametangium (Pl. XI, fig. 42). One or more very large oil drops and much granular material are always visible in the interior.

All attempts to obtain germination have failed, although zygosporae have been set up under many conditions, and about two hundred have been kept for over two years. Light, darkness, heat, freezing, dissolving the walls with acid, squeezing off the outer coat, and weak and strong liquid and solid media have been tried without success.

(4) *Abnormal zygosporae*

The suspensor of the large gametangium frequently exhibits one or more incomplete annular septa, similar to that which cuts off the gametangium (Pl. XI, fig. 35). In the abnormal zygosporae shown in Text-fig. 80, such a septum forms the boundary of the spore, and it is to be presumed that the second septum becomes complete before the first. More often the whole female sac becomes thick-walled (Text-fig. 79, and Pl. XI, fig. 45). Up to 2 % of these zygosporae occur in some cultures, and so far as can be seen, the first gametangium wall is never complete, and the outer walls are always thinner than in the normal zygosporae. Such abnormal zygosporae appear to be unique in the Mucorales. The only forms which resemble them are the thick-walled spores described by Chesters⁽¹¹⁾ in his recently discovered *Azygozygum chlamydosporum*. These, which he interprets as imperfect zygosporae, have, however, a different origin, as they consist of the two gametangia pressed together, with or without perforations in the wall between them.

I have seen no azygosporae, nor chlamydosporae, in any of the cultures of *Dicranophora* grown over a period of two years. *

(5) *Comparison with previous accounts*

I am in disagreement with the following statements made by previous authors:

That the large gametangium always arises from an outgrowth formed in response to contact with the male hypha (Blakeslee and Ling-Young).

That the two gametangia are completely isolated before the wall between them breaks down (Ling-Young).

That the zygosporae is spherical (so described by Schröter and not denied by the others).

In addition the suspensors have been observed to branch, but not to give rise to sex organs, as reported by Ling-Young.

The fact that the sexual branches are not necessarily in contact from the first is implied by Ling-Young's reference to "l'action provocatrice du gamétophore mâle, qui évolue plus rapidement", but is nowhere stated or illustrated by his figures. In this respect *Dicranophora* alone of the Zygomycetes appears to resemble *Phycomyces nitens* as described by Burgeff⁽¹²⁾.

The "characteristic bulge" which "develops" on the stalk of the larger gametangium, as described by Gwynne-Vaughan and Barnes, is almost certainly an unfertilized female lobe. Where, as frequently happens, only one of a group of female sacs is fertilized, the others appear to be outgrowths from its stalk; sometimes, however, the fertilized sac is itself lobed or branched at the base; but in either case the bulge is present before the gametangium is formed.

The "smoothness" attributed to the zygospore by Schröter is probably, as suggested by Vuillemin, due to his having examined young zygospores on which the thin gametangial membrane was still present.

DISCUSSION

(1) *Sexuality*

Dicranophora possesses a character which is unique in the Mucorales, namely, the large female organ, which is clearly recognizable whether it is fertilized or not. The size difference between the gametangia is invariable, the female sac being never less than three times the diameter of the male.

Heterogamy is a much commoner phenomenon in the Mucorales than was previously thought, but the size differences reported in many genera, e.g. *Rhizopus* (Blakeslee⁽¹³⁾), *Sporodinia* (Moreau⁽¹⁴⁾), and *Mucor hiemalis* and *Absidia glauca* (Ling-Young⁽³⁾, p. 493), are all very variable, and are regarded as having a nutritional rather than a sexual significance. A few species only are reputed to show pronounced and constant heterogamy, viz. *Dicranophora fulva*, *Zygorhynchus* (several species), *Absidia spinosa*, and species of *Syncephalis* (especially *S. nodosa*).

Of these, Green⁽¹⁵⁾ has shown that in *Zygorhynchus* the size difference is far from constant, and may even be reversed. My observations on *Absidia spinosa* show that here also the relative sizes are very variable, the only constant difference being the growth of appendages on one suspensor only. The fact, referred to in a recent paper by Dodge⁽¹⁶⁾, that when *Zygorhynchus* and *Absidia spinosa* are grown together "large cell reacts with large cell, small with small" indicates that in these forms the inconstant size difference does not correspond with any functional sex difference. On the other hand, in *Syncephalis* (Thaxter⁽¹⁷⁾) where the functional difference is marked, the heterogamy does not consist in a size difference, but the sex organs resemble those of the

Plectomycetes, and are entirely different from those of *Dicranophora*, which appears to be the only member of the Mucorales in which a constant size difference is established. The appearance of the young sexual branches in *Dicranophora* (e.g. Pl. XI, fig. 28) immediately suggests an antheridium and oogonium, and makes a comparison with the sexual stage of the Oomycetes inevitable.

Points of similarity are as follows:

(1) The female organ may have an independent existence apart from sexual contact.

(2) The female organ is clearly distinguishable from the male in size and shape, and contains a very large amount of reserve material.

(3) The whole contents of the male are discharged into the female sac.

(4) The protoplast of the zygote is entirely contained within the female organ.

These differences are clearly sexual, in the narrow sense of the term employed by Blackman (18), although the observation of Ling-Young, which I have been unable to confirm, that in *Dicranophora* both large and small gametangia can give rise to hyphae bearing both types of sex organ, indicates that there is here no segregation of nuclei of opposite sexes. *Dicranophora*, however, is the only Zygomycete which shows pronounced and invariable oogamy, and the similarity of its sexual stage to that of such an Oomycete as *Albugo Bliti*, in which multiple fusion takes place, can scarcely be ignored. It possesses, nevertheless, the characteristics of a very specialized Zygomycete, and even the sexual stage differs from that of the Oomycetes in the following important points:

(1) The female branch in *Dicranophora* is capable of indefinite growth and reversion to the vegetative state.

(2) The "antheridium" is included in the zygote as a small boss on the zygospore wall.

(3) The "oogonium" is not completely isolated until after the entry of the contents of the "antheridium".

(4) The nuclear phenomena of the sexual stage, as reported by Ling-Young (3), are essentially similar to those of a homogamic Zygomycete such as *Sporodinia grandis*.

(2) Systematic position

The wide range of forms shown by *Dicranophora*, each bearing some resemblance to a form shown by another genus of the Mucorales, is one of the peculiarities of the fungus. Instances are the *Pilobolus*-like swellings, and oedocephaloid heads occasionally produced on the sporangial stage, but a comparison with the four genera to which it shows the greatest resemblances is most concisely shown by a table:

Comparison of other Mucorales with Dicranophora fulva

Genus	Similarities	Differences
<i>Thamnidium</i>	Large and small sporangia, the latter on dichotomous branch systems; apparently homothallic.	No pigment, sporangioles without columella, shed as conidia. Zygo-spores homogamic where known, in scalariform series on erect filaments. Usually coprophilous.
<i>Zygorhynchus</i>	Homothallic, usually markedly heterogamic. Copulation by large curved branch commonly occurs as in <i>Dicranophora</i> (Text-fig. 69). Zygosporangia non-aerial.	No pigment. Sporangia dissimilar. Heterogamy variable. Copulation unlike that usual in <i>Dicranophora</i> . Zygosporangia dissimilar. Not epiphytic.
<i>Sporodinia</i>	Coarse hyphae with granular pigmented contents. Sporangia on dichotomous branch systems. Homothallic, occasionally heterogamic. Zygosporangia also dichotomous. Parasitic on wide range of Agarics and Boleti.	No sporangioles. Gametangia normally about equal, and copulating branches straight. Zygosporangia aerial.
<i>Spinellus</i>	Pigmented hyphae. Spindle-shaped sporangioles. Homothallic. Occasionally heterogamic (<i>S. chalybeus</i>). "Tong-like" copulation. Crevised zygosporangia coat. Parasitic, each species restricted to one species of Agaric.	"Thorny" hyphae. No sporangioles. No prominent dichotomous branch systems. Normally homogamic. Zygosporangia aerial.

A consideration of the facts summarized in the table leads inevitably to the support of Vuillemin's view, that *Dicranophora* should be classified with *Sporodinia* and *Spinellus*. My investigation has, I think, shown that Vuillemin was wrong in supposing that the sporangioles lacked a columella, and could be shed, like those of *Thamnidium*, and it is clear that they are quite comparable with the sporangia of *Sporodinia*, except in size and spore number. With regard to its range of host also, *Dicranophora* seems to be intermediate between *Sporodinia* and *Spinellus*, since it seems to be confined to Boleti and the lower Agarics, although the number of times it has been found is too small to make this assumption reliable.

In any phylogenetic series, it is convenient to regard this group of homothallic Mucorales (*Dicranophora*, *Sporodinia*, and *Spinellus*) as linked to some extent to the Oomycetes, through the oogamy of *Dicranophora*, the Cephalidaceae (especially *Syncephalis*) being regarded in a similar way as a link with the Ascomycetes. This, of course, is not a suggestion that *Dicranophora* is either derived from, or has given rise to, any Oomycete, which, in view of its specialized habitat and asexual characters, is highly improbable; but our objective knowledge of the phylogeny of the Phycomycetes is so negligible, that the chief justification for any such phylogenetic arrangement, based upon comparative morphology, seems to me to lie in its value as a mnemonic for the characters of the fungi.

SUMMARY

1. *Dicranophora fulva* grows readily in culture, but forms no zygospores on purely artificial media. It is extremely sensitive to light, forming sporangia only when illuminated, and zygospores only in the dark, or in very diffuse daylight.

2. In the material studied, the columella in the large sporangium is rounded, not conical. There is a columella in the sporangiole, but the claw-like outgrowths described by previous authors have been observed only once. The columella in both large and small sporangia has, however, been seen to produce a hypha.

3. The reported dimorphism of the spores in the two sorts of sporangia does not exist, except in so far as the spores in the sporangiole are somewhat more compressed. The liberation of the contents of a large sporangium as a single giant spore is described. The range of spore size appears to be far greater than in any other Mucorales.

4. The sporangiole differs from the sporangium only in its size and position on a dichotomous branch system. Intermediate forms occur which are not distinguishable as one or the other.

5. The sporangiophore appears to give rise to about the same number of spores, whether it bears one large sporangium, or forms a branch system bearing many sporangioles.

6. *Dicranophora* is the only Zygomycete in which a constant size difference between the gametangia has been shown. It is unique in possessing a female organ which may have an independent existence apart from sexual contact, and it differs from most of the Mucorales in that the sexual branches grow into contact, and are not necessarily in contact from the first.

7. The extent to which the female branch produces an outgrowth towards the male is very variable. The wall between the gametangia breaks down before the larger one is completely isolated, and the whole contents of the male pass into the female gametangium, in which the zygospore is formed.

8. The zygospore is not spherical, but slightly flattened on one side, with a deeply creviced outer coat, and a small boss representing the male gametangium. It has not been germinated. Some abnormal zygospores are described.

9. Support is given to the view that *Dicranophora* should be classified with *Sporodinia* and *Spinellus*, rather than with *Thamnidium*. Certain marked resemblances to the Oomycetes are also pointed out.

In conclusion I wish to thank Prof. R. R. Gates, at whose suggestion the work was carried out, for his advice and criticism, and also Mr C. S. Semmens for developing and printing the photo-micrographs.

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EXPLANATION OF PLATES X AND XI

PLATE X

Dicranophora fulva Schröt. The sporangial stage

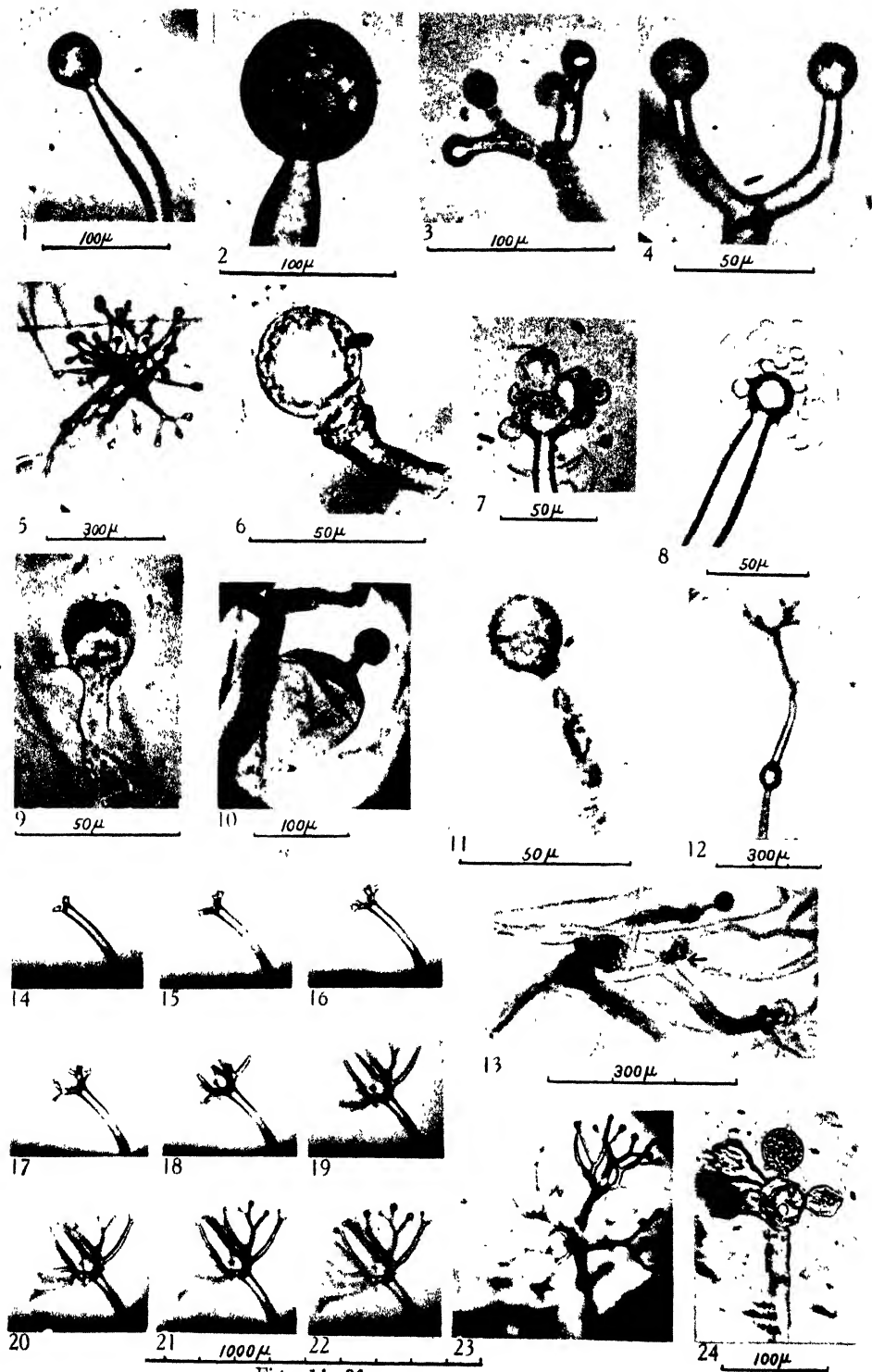
- Fig. 1. Sporangium showing columella.
- Fig. 2. "Sporangial drop" showing shape of columella and spores.
- Fig. 3. Part of sporangiole branch system.
- Fig. 4. Sporangioles showing early stage in formation of columella.
- Fig. 5. Sporangiole branch system.
- Fig. 6. Single "giant" spore (30 μ in diameter) in contact with a columella which shows traces of the broken membrane.
- Fig. 7. Burst sporangium having a small number of spores.
- Fig. 8. Patch of spores formed by the bursting of a sporangium of average size against the cover-slip of a cell.
- Fig. 9. Sporangiole mounted in water, showing columella, membrane, and two spores. Taken with $1/12$ in. objective.

- Fig. 10. Large subsporangial vesicle, separated from the sporangium by a short stalk. Mounted material.
- Fig. 11. Sporangiole showing surface encrusted with crystals.
- Fig. 12. Sporangial swelling which has produced a branch bearing sporangioles.
- Fig. 13. Branch produced by the columella of a burst sporangium, the position of which is indicated by the arrow. The branch has begun to form a young sporangium, but has burst owing to immersion in water. The apparent "collar" round the base of the young sporangium is extruded contents.
- Figs. 14-23. A series showing the development of a sporangiole branch system. Times, in hours and minutes from the first photograph: Fig. 14 (0); Fig. 15 (0.30); Fig. 16 (1.0); Fig. 17 (1.30); Fig. 18 (3.20); Fig. 19 (4.50); Fig. 20 (6.50); Fig. 21 (7.50); Fig. 22 (8.30); Fig. 23 (20.0). The last few figures are partly obliterated by a creeping film of water.
- Fig. 24. "Oedocephaloid" head bearing five radially arranged "buds" some of which are sprouting *in situ*—produced by a sporangiophore in a moist culture.

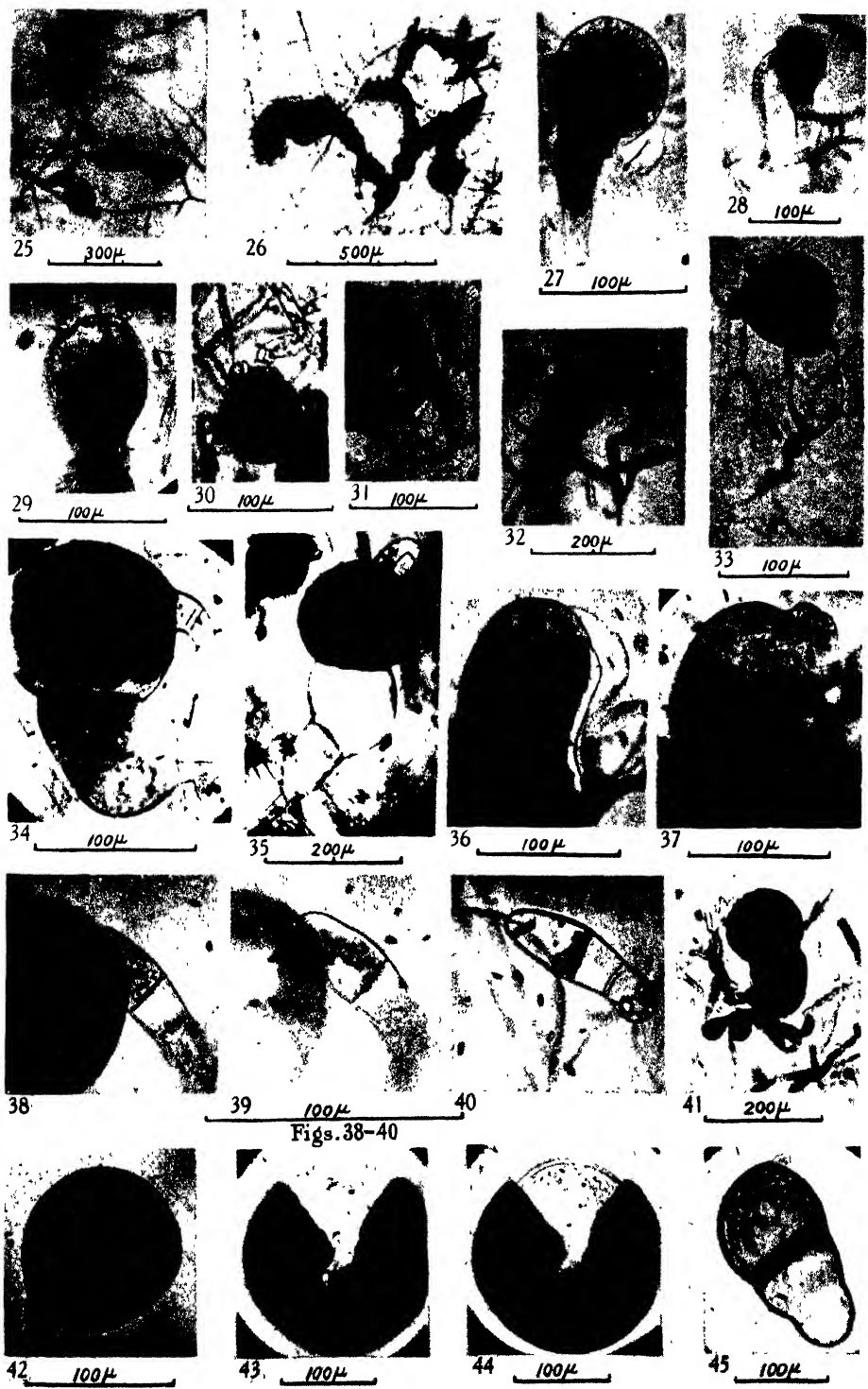
PLATE XI

The sexual stage

- Fig. 25. Group of female "sacs".
- Fig. 26. Large female zygophoric branch system, bearing two zygospores, and showing dichotomous branching.
- Fig. 27. Single female sex organ, showing granular contents.
- Fig. 28. Very early stage in conjugation, probably before contact.
- Fig. 29. Early contact. Formation of slight papilla on female branch.
- Fig. 30. Later. Considerable outgrowth of female towards male.
- Fig. 31. Formation of septum on the male.
- Fig. 32. Beginning of female septum. Male gametangium relatively clear of granular contents.
- Fig. 33. Young zygospore, showing "antheridial boss" and remains of protoplasm in the large suspensor.
- Fig. 34. Earlier stage than Fig. 33, showing thickening of "antheridial" wall.
- Fig. 35. Young zygospore with terminal "antheridium" and incompletely septate large suspensor.
- Fig. 36. Very early contact of the two sex organs.
- Fig. 37. The same, 13½ hours later; the shadows are due to water drops, and encroaching hyphae.
- Figs. 38, 39, 40. Stages in the passage of contents of a male into a female gametangium which was losing its contents owing to the effects of illumination.
- Fig. 41. Developing zygophoric sac, with a group of unfertilized lobes at its base.
- Fig. 42. Zygospore in optical section, showing large oil drop, and the complete obliteration of the lumen of the "antheridium" by the thick double wall.
- Fig. 43. Crushed zygospore, focused to show the crevices in the outer coat.
- Fig. 44. The same, focused to show the inner hyaline wall, and contents.
- Fig. 45. Abnormal zygospore, with imperfect gametangial septum.



Figs. 14-23



NOTE ON A RARE BEETLE, *CARTODERE FILUM* AUBÉ, EATING FUNGUS SPORES

By H. D. GORDON, B.Sc., Ph.D.

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(With Plate XII)

FOR a number of years it has been noticed that the spores of certain fungi kept in a dry condition in the laboratory of the Royal Botanic Garden, Edinburgh, become compacted, after a time, into little cylindrical masses, which strongly suggest that the spores have been ingested by some small animal, and egested in the faeces. The fungi attacked in this way are those having dry, powdery spores, e.g. species of *Lycoperdon*, *Ustilago* and *Tilletia*.

The faecal pellets (Pl. XII, fig. 1) are roughly cylindrical, straight or slightly curved, generally appearing under the microscope as oblong masses the ends of which may be clean-cut, or more or less irregular. The diameter ranges from 23–37 μ , but much wider variations occur in the length of the masses, measurements from 40 to 200 μ having been obtained; this is readily understandable, as the diameter must be conditioned by the dimensions of the alimentary canal of the animal depositing them, whereas the length of the individual fragments breaking off on deposition is largely fortuitous. These pellets seem to be made up entirely of closely compacted spores, which retain their normal appearance. Presumably they bear a thin coating of mucid substance, to which their cohesion is due, but no interstitial matter is discernible under the microscope. The number of spores which may be counted in the transverse diameter of a pellet, e.g. at one end, depends on the spore size of the species concerned, as illustrated by the following average figures relating to the three species most noticeably affected in the laboratory:

	Average spore size μ	Average number across pellet	Average size \times average number μ
<i>Lycoperdon pyriforme</i>	3.9	7	27.3
<i>Ustilago Avenae</i>	5.7	5	28.5
<i>Tilletia Tritici</i>	18.6	2	37.2

The figures for number of spores across the pellet must be taken as approximate only, for the spores are not found arranged in definite rows; being roughly spherical bodies they tend to alternate with each

other. The relatively large spores of *Tilletia Tritici* are never arranged more than two deep (Pl. XII, fig. 2), and actually they are usually a little out of direct alignment except with spores under average size, so that the diameter of the pellets is somewhat less than the product shown in the final column of the table. Making allowance for this, it will be noted that the figures fall within the range already quoted for the diameter of the pellets. It is evident that this somewhat elastic limit is independent of the spore size, and is imposed by the agent responsible for the formation of the pellets, as we should expect if this is the alimentary canal of a small animal.

For some time no animal which might be responsible for this transformation of the spores was noticed, and its presence was merely inferred from the observed effects. Recently the appearance of faecal pellets in the spores of some smutted oats which had been in store only a few months afforded the opportunity for a systematic search, and a small brown beetle was found in considerable numbers. Further examination proved that this was responsible for the damage.

Specimens of the beetle were submitted to Dr A. E. Cameron, of the Edinburgh University Department of Entomology, who identified the beetle as *Cartodere filum* Aubé, of the family Lathridiidae (Clavicornia), and kindly supplied some further information about it. Fowler (1889) comments on this species as follows:

"Very rare; it appears to be chiefly confined to herbaria, although it occasionally occurs in fungi in other countries. Burton-on-Trent (Mr Mason's herbarium, in some small numbers); Scotland, Edinburgh (found by Prof. McNab in the herbarium of the Royal Botanic Gardens)."

The beetle has evidently found, amongst the dried specimens of various fungi, conditions of life and food material which suit it admirably, with the result that it has survived for many years, and now appears to be plentiful and flourishing.

Specimens of *Ustilago Avenae* and *U. Hordei* collected in Wales in August 1935 and stored in the affected cupboards showed no sign of damage in October of the same year, but when these specimens were again being used in February 1936, faecal pellets were abundant in the smutted ears. It was amongst these specimens that search was carried out, and many living mature beetles were found, as well as a fair number of larvae and a few pupae. The beetle was seen to deposit the pellets quite rapidly.

A few beetles were killed, dehydrated in absolute alcohol, cleared in clove oil and mounted in Canada balsam. The contents of the alimentary canal could now be clearly seen, apparently consisting entirely of smut spores which, in the hinder gut, were compacted together into a continuous cylindrical mass (Pl. XII, fig. 3), which broke up on emergence into the characteristic pellets.

The larvae were observed to deposit similar, but generally somewhat more slender pellets, and when a few larvae were cleared and mounted, the spores could be seen in the gut (Pl. XII, fig. 4). They were also present in the gut of the pupae, though these, being quiescent, did not deposit pellets. Fowler (1889) says that "the larvae [of Lathridiidae generally] probably feed on cryptogamic substances, the excrement and skin of various insects, etc." Here larvae and adults alike were apparently subsisting on the same pure diet of one cryptogamic substance, namely smut spores.

The unaffected spores of *Ustilago Avenae* and *U. Hordei* (collected August 1935) were still viable in February 1936, germination beginning after a few hours in water. As the spores constituting the faecal pellets were not visibly altered except as to their arrangement, these were tested for viability. To ensure that the spores used had actually passed through the alimentary canal, a living beetle was kept under observation in a clean hollow-ground slide, where it was seen to deposit pellets of spores. These were transferred to a hanging drop of water.

Most of the spores failed to germinate, but a few produced their promycelia and sporidia within eighteen hours, when the photograph (Pl. XII, fig. 5) was taken; by this time the bacteria present in the faeces had multiplied very considerably. The relatively long period of eighteen hours should not be stressed too heavily; the spores had germinated overnight within this time, and even in experiments with unaffected spores some always took a good deal longer than others to germinate. But one evident difference was the very small number of spores which had germinated even after eighteen hours. In drops where both unaffected spores and pellets were present the promycelia of the former were generally so abundant as to hamper observation of the latter, and frequently when a promycelium appeared to proceed from one of the pellets the possibility remained that it might belong to an unaffected spore adhering to the mass. But where fresh pellets only were introduced into the drop, it was very noticeable that most of the spores showed no signs of germination.

From this it appears that, while spores may survive and germinate after passage through the alimentary canal of the beetle, they are nevertheless subject to some adverse influence to which most of them succumb. This lethal effect was probably caused by the digestive juices of the beetle.

The exact composition of the spore wall of *Ustilago* appears to be unknown, but it may be taken that it consists mainly or entirely of some form of cellulose. Microchemical tests show that it is not the same form of cellulose as is found in the cell walls of the higher plants and of a few fungi, such as *Peronospora*. For example, spores of *Ustilago* and *Peronospora* were mounted together and tested with

sulphuric acid and iodine, when the spore wall of *Peronospora* gave the blue colour reaction characteristic of cellulose, while that of *Ustilago* simply assumed a yellowish brown tint. It has been customary to refer to the substance of such walls as "fungus cellulose", as it is believed to consist essentially of cellulose, though in a slightly different form from that occurring in the majority of plants.

If this be so, it would appear from the reduced germination capacity of the egested spores that these must have been acted upon by an enzyme capable of dissolving, at least in part, their protective covering, that is to say, by a cellulase.

According to Mansour & Mansour-Bek (1934) no evidence of the presence of cellulase in insects had been obtained prior to 1919, and any digestion of cellulose which might take place was believed to be due to the presence of micro-organisms in the intestinal tract. Subsequent to 1919, however, cellulase has been recorded in a number of insects, including wood-eating beetles. It is therefore quite possible that *Cartodere filum* may secrete cellulase, especially as the herbarium material on which it generally lives must contain a high percentage of cellulose, but I know of no experimental information on this point.

Reference may be made here to the digestion of spores of *Tilletia Tritici* by animals. These spores have been found in the faeces of several vertebrates (man, dog, rabbit, guinea-pig, etc.) and appear to be affected but little if at all by their passage through the digestive tract (Dobson, 1926). Conflicting results have been reported as to their capacity for germination when obtained from the faeces. In vertebrates the breakdown of cellulose is generally admitted not to be the work of a cytase, but due mainly to the action of intestinal bacteria. It is possible that the conflicting results may be related to variations in the composition of the bacterial flora.

It was not possible to test the viability of egested *Tilletia* spores, as the material available was old, and even the unaffected spores were not capable of germination. The spores of *Ustilago* on which the observations of germination were made were six months old; fresher spores might have germinated more freely.

No attempt has been made here to deal in detail with the entomological aspect of the problem. The purpose of this note is rather to record the agent responsible for the appearance of these characteristic pellets amongst dry fungus spores, the persistence in the laboratory of this rare beetle, and the fact that spores are capable of germinating after passing through the alimentary canal of the beetle, but do suffer a reduction in their capacity for germination.

I am indebted to Dr Malcolm Wilson for the suggestion that this problem might be worthy of investigation, and its explanation of interest to other workers.



Fig. 1

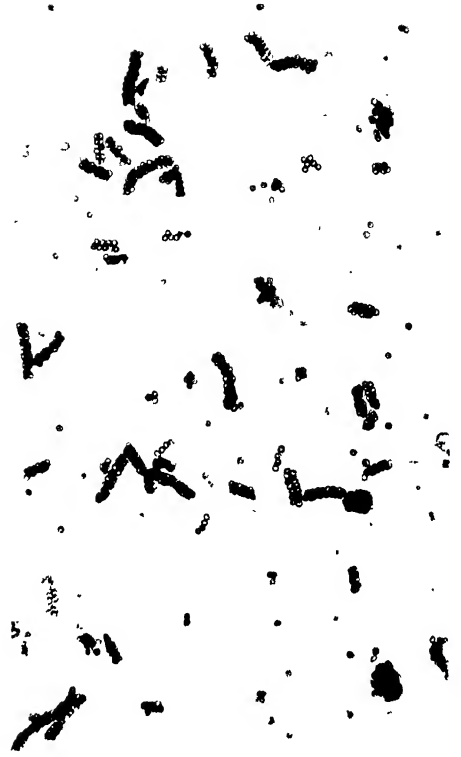


Fig. 2



Fig. 3



Fig. 4



Fig. 5

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EXPLANATION OF PLATE XII

- Fig. 1. Basidiospores of *Lycoperdon pyriforme*, egested by *Cartodere filum*, as seen under a low power.
- Fig. 2. Chlamydospores of *Tilletia Tritici*, egested by *Cartodere filum*.
- Fig. 3. *Cartodere filum* Aubé; the mature beetle cleared to show the dark mass of spores of *Ustilago Avenae* in the gut.
- Fig. 4. *Cartodere filum*; larva with spores of *Ustilago Avenae* in the gut.
- Fig. 5. Chlamydospores of *Ustilago Avenae*, egested by *Cartodere filum*; photographed after eighteen hours in water; a few spores in the smaller mass have germinated.

CONTRIBUTIONS TO THE STUDY OF *PENICILLIUM EGYPTIACUM* VAN BEYMA

By YOUNIS S. SABET

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(With Plate XIII and 10 Text-figures)

INTRODUCTION

THE genus *Penicillium* includes several hundred species, and is cosmopolitan in distribution. Most species, so far as is known, multiply exclusively by means of conidia, but in a few species perithecia may develop. This development may depend upon some special cultural or other environmental condition, such as diminution of the supply of free oxygen(3, 21) or an abundant supply of sucrose(1). Heterothallic(6) and homothallic(18) species have been described. Several authors(7, 12, 13, 15, 20) have found perithecia in species of *Penicillium* without ascribing their formation to any special circumstances.

P. egyptiacum, a form isolated from soil in Egypt(16), develops perithecia in great numbers on ordinary media(17). It is proposed to give some account of the morphology of this species, and of its behaviour in culture.

EXPERIMENTAL WORK

Cultural characters

P. egyptiacum has been grown on more than thirty media of very different nutritive values. These media included synthetic agars, plant decoction agars, and nutrient solutions, as well as plugs of vegetable material of divers kinds. The more striking results of these experiments are summarized below.

On Conn's medium and Barnes's medium, the mycelium formed thin, smooth, shining and somewhat transparent mats. Aerial mycelium developed as rope-like strands of inter-twisted hyphae bearing conidiophores. The perithecia were scattered over the surface of the medium in fair numbers (Pl. XIII, fig. 1), but they were not crowded together.

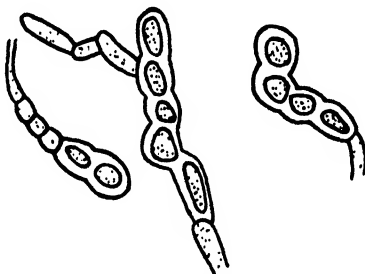
On malt agar, date extract agar(16), Brown's agar, and Richard's solution agar, the comparatively short and crowded hyphae produced a dense growth. Aerial mycelium was luxuriant. Young colonies were white, not only because of the closely packed hyphae, but because of the enormous numbers of perithecial primordia. As the

latter matured, they imparted a brown, granular aspect to the colony. On Richard's solution agar, the mycelium and the substratum became wrinkled, and later, cracked (Pl. XIII, fig. 2).

Some preliminary tests were made on the effect of the source of nitrogen. For this purpose, a medium of the following general composition was prepared:

	%
Glucose	2
Tripotassium phosphate	0.5
Magnesium sulphate	0.17
Agar	2
A nitrogenous compound	1

When ammonium chloride was used, conidiophores developed in large numbers, and the chains of conidia were exceptionally long. The colonies were velvety in texture and uniformly green, owing to the abundant conidia, but perithecia were rare. When asparagin supplied the nitrogen, growth was soon arrested by staling, and only very few perithecia appeared.



Text-fig. 1. Oidia-like bodies developed on lime fruit. $\times 550$.

Growth was weak, and perithecia scanty, on a medium containing filter paper as the only source of carbon.

Strong vegetative development, with heavy crops of perithecia, was obtained on plugs of potato and of carrot. *P. egyptiacum* grew and fruited well on the fruits of orange, mandarine and lime; on the latter, the mycelium showed a special tendency to produce thick-walled structures resembling oidia (Text-fig. 1). The addition of compounds of arsenic to media led to the formation of short, richly branched hyphae; these cultures did not emit a garlic-like odour.

Saltation

Sectoring was frequently observed; in general, the sectors consisted either of a dense, brown, granular growth rich in perithecia, or of a green velvety area, covered with conidiophores, but containing very few perithecia. A transfer to malt agar from a green sector gave a homogeneous green colony which retained these characteristics

through the four subsequent transfers that were made (Pl. XIII, figs. 3, 4).

Zonation

Zonation was observed occasionally on Brown's agar and malt extract agar, and also when cultures were grown under reduced atmospheric pressure. The saltant which produced heavy crops of conidia showed zonation on malt agar and on Barnes's medium.

Influence of temperature

Three cultures were grown on malt agar at each of the following temperatures: at about 4° C. in an ice-box, at 13–14·5, 18, 22, 26, 30 and 34° C. The results appear in Table I.

Table I. *Influence of temperature on Penicillium egyptiacum*

Days	Diameters (in mm.) of colonies at						
	4° C.	13-14·5° C.	18° C.	22° C.	26° C.	30° C.	34° C.
2	3	6	6·7	7·3	7·3	7·5	3
4	No more growth		10·2	13·3	13	12	5
6	„	15	17·7	18·2	18	15·4	No more growth
8	„	19·5	22·7	23	23·5	20·4	„
10	„	24	26·7	28	28	23·5	„
12	„	26	29	32	32	28	„
14	„	30	33	36	36·5	31	„
30	„	—	60	69	72	52	„

The diameters are the averages of two measurements taken from each culture of the triplicate sets.

These data indicate that the optimum temperature for growth lies between 22° and 26° C. There is, however, little difference in the strength of growth at any temperature between 14° and 30° C. In all cultures where growth continued, perithecia developed in the normal manner.

Some cultures were left for six weeks at 4° and at 34° C., transfers being then made to malt agar slants, which were incubated at 22° C. Since normal colonies resulted, it is evident that the fungus had suffered no injury during the time it had been exposed to temperatures unfavourable to growth.

Influence of the pH of the medium

Penicillium egyptiacum was grown on Richard's solution agar, adjusted by additions of phosphoric acid or of sodium bicarbonate to various hydrogen-ion concentrations. After autoclaving, the pH of the agars was determined by the colorimetric method (4). From the many media prepared, samples were selected with the following range of pH: 3·8, 4·8, 5·8, 6·8, 7·8 and 8·8. Three Petri dishes were poured

from each lot of these media, inoculated, and incubated at 22° C. The results are summarized in Table II.

Table II. *Influence of pH of the medium on Penicillium egyptiacum*

Days	Diameters (in mm.) of colonies at pH					
	3·8	4·8	5·8	6·8	7·8	8·8
2	6	6	4·25	4·5	4	4·5
4	18	16	15·75	14·75	14	13·75
6	28·75	24	27	25·5	22·5	26
8	36	31·25	36	35	30·5	33
10	45·75	38·75	43	43	38·5	39
20	78	73	76	77	75	75

The diameters are the averages of two measurements taken from each culture of the triplicate sets.

The figures in Table II indicate, that within the range tested, the pH of the medium has no significant effect upon the growth of *P. egyptiacum*. It was noted, however, that perithecia developed later on the media with a pH 8·8 than on the other media.

Influence of atmospheric humidity

Six large crystallizing dishes were sterilized and provided with a range of aqueous solutions of calcium chloride⁽¹⁴⁾ giving the following atmospheric humidities at 20° C.: 100, 96, 90, 80, 70 and 60 %.

Twelve small Petri dishes containing malt agar were inoculated with *P. egyptiacum* and attached in pairs, upside down, to the inside of the lids of the crystallizing dishes. The lids of the Petri dishes were removed so that the media were freely exposed to the atmosphere in the larger dishes. Access of air was checked as far as possible by packing cotton-wool into the spaces between the lids and bottoms of the crystallizing dishes.

The preparations were incubated at 20° C. for ten days, when the diameters of the colonies were measured. The results appear in Table III.

Table III. *Influence of atmospheric humidity on Penicillium egyptiacum*

Moisture %	Diameter of colony mm.	Characters of colony
100	26	Powdery brown, perithecia crowded
96	25	" " " "
90	14	Velvety, greenish, perithecia scanty
80	9	Velvety, greenish, perithecia absent
70	6	" " " "
60	5	" " " "

The diameters are the averages of two measurements taken from each culture of the duplicate sets.

It seems that the formation of perithecia occurs freely only in very moist atmospheres. At about 90 % humidity, conidial formation assumes the upper hand, and perithecia become rare, while at 80 % humidity and below, perithecia cease to form. These results are in agreement with those obtained by Blochwitz⁽²⁾ and by Schwartz⁽¹⁹⁾ with species of *Aspergillus*.

Influence of light

Six dishes of malt agar were inoculated with *Penicillium egyptiacum*. Three were covered with black enamel and placed in a large dish, similarly darkened. The remaining three dishes were placed in a large, undarkened dish. Both preparations were left for thirty days on a bench opposite to a window. It was then found that the colonies grown in the light averaged 70 mm. in diameter, while those grown in the dark averaged 45 mm. in diameter. All the cultures had produced perithecia. Those developed in the light were scattered, and less numerous per unit area than the perithecia in the darkened dishes. It appears therefore that light favours the mycelial growth of *P. egyptiacum* and depresses the formation of perithecia, darkness having the opposite effect.

Influence of atmospheric pressure

Two batches of malt agar slants were inoculated with *P. egyptiacum* and incubated at 25° C. for twenty-four hours. When it was clear that growth had begun, one batch was placed in a desiccator attached to an air pump. This was run for three hours, when the tap of the desiccator was closed. The other batch was placed in a desiccator with the tap open. Both were left for a month at 18–22° C. The tap of the exhausted desiccator was then opened, and since air was heard to rush in, it was evident that the fungus had been under reduced pressure during the whole time of the experiment. The cultures grown under reduced pressure were zoned with alternate bands of greenish conidial areas and closely compact, brown ridges of perithecia; the cultures kept at atmospheric pressure were even and unzoned.

The morphology and cytology of Penicillium egyptiacum methods

Slides were thoroughly cleaned from grease and other dirt, placed in Petri dishes, sterilized, and covered with a thin layer of Barnes's medium. This medium was used, since it is clear, and being of low nutritive value, the mycelium grows loosely on it, and the scattered perithecial rudiments are easy to examine. The cultures were inoculated and incubated at 25° C., for four or five days. The fungus was then fixed by adding to the dishes a quantity of strong Flemming's

solution diluted with an equal volume of water. After washing, and taking up through the usual series of alcohols, the material was stained by the glycerine-erythrosin method⁽¹¹⁾. Up to this point, all operations were done on the fungus in the dish, care being taken to disturb it as little as possible. The slides, covered by a thin film of agar, were then removed, and the stained material mounted in warm glycerine jelly. The preparations were sufficiently thin to be examined with an oil immersion objective.

Other material, grown and fixed in similar manner, was stained in Heidenhain's iron haematoxylin, and used for a study of the nuclei in the conidiophores and conidia.

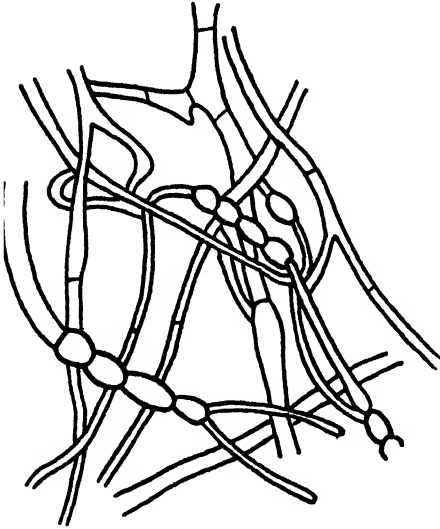
In order to investigate the perithecia, the fungus was grown on malt agar, fixed in the fluid 2BD of La Cour⁽⁵⁾, embedded in paraffin, cut at 5μ , and stained with iron haematoxylin, with erythrosin in clove oil as a counter-stain.

Development of the reproductive organs.

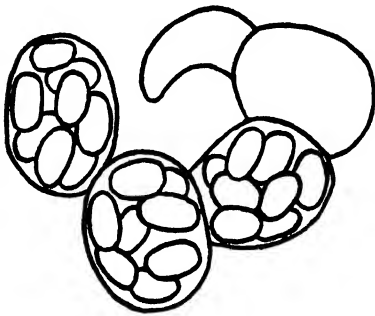
Perithecia. Some hyphae rise slightly above the substratum and form extensive branch systems with frequent anastomoses. They give rise to aerial knots, which are the primordia of the perithecia. Certain stout hyphae, distinguished by their close septation and rich contents, become centres of activity and are soon surrounded by wefts of anastomosing hyphae. These hyphal masses are soon surrounded by a pseudo-parenchymatous envelope, and within this the asci subsequently develop. Sometimes complicated hyphal masses are found in which the stout, septate hypha cannot be clearly distinguished. Spirally wound primordia have never been seen, nor has anything suggesting an antheridium been found.

Serial sections of young perithecia reveal a mass of solid ground tissue with the remains of darkly stained septate hyphae embedded in its centre (Pl. XIII, fig. 5). These hyphae are still distinguished by their rich contents and thin walls, and although they may be somewhat curved, they are not spirally wound. It seems the archicarp of *P. egyptiacum* is merely a stout, closely septate hypha (Text-fig. 2), rich in contents, and but slightly curved. It puts out branches, closely septate and rich in contents, and these give rise to much narrower ascogenous hyphae, which are short and curved in a characteristic fashion (Text-fig. 3). Each becomes septate, and each segment then becomes an ascus (Text-fig. 5c, d). The first branches to emerge from the archicarp are multinucleate, as are the young ascogenous hyphae which contain pairs of nuclei in close association. The septa form between the pairs of nuclei, cutting off the rudiments of the asci. The first ascus matures at the tip of the ascogenous hypha, and a second, third, and sometimes a fourth (Text-fig. 4) ripen in succession below it; the asci are thus formed in chains.

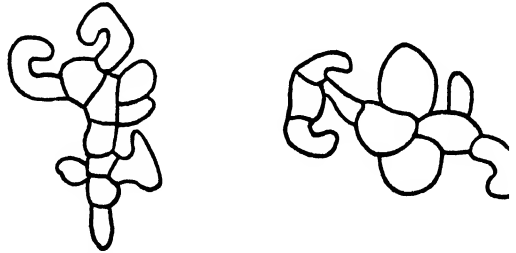
Young asci containing four nuclei were recognized, but as nuclear fusions have not been detected, it is not clear if all the binucleate cells that were observed were of the same kind. Some were doubtless the binucleate segments of the ascogenous hyphae, others were probably young asci in which the first nuclear division had occurred. It seems very probable that the two nuclei fuse in the young ascus, and that



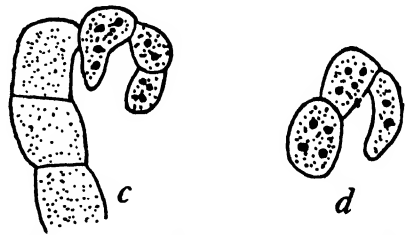
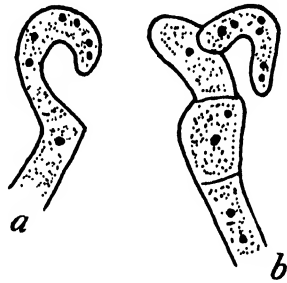
Text-fig. 2. Perithecial initial. $\times 666$.



Text-fig. 4. Asci drawn from fresh perithecia. $\times 2500$.



Text-fig. 3. Ascogenous hyphae drawn from young fresh perithecia. $\times 1500$.

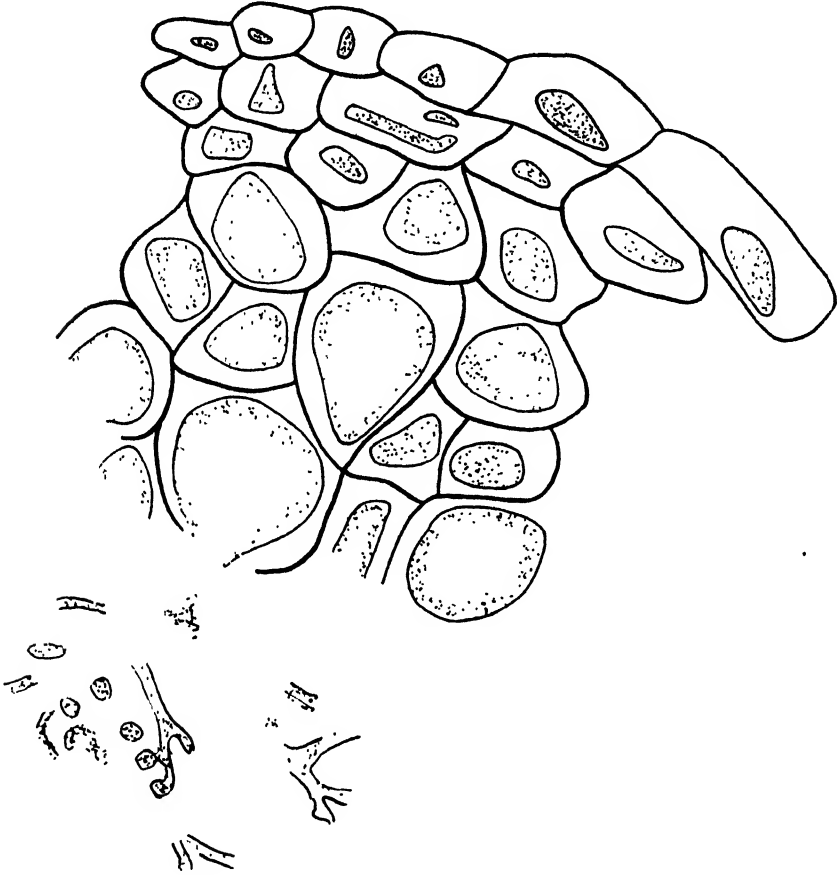


Text-fig. 5. Ascogenous hyphae in different stages of development. $\times 2500$.

three rapidly succeeding divisions give the eight nuclei for the eight ascospores. These may be seen clearly in the ascus (Text-fig. 4), as Emmons has already shown (8).

The outer layers of the fruit body are compact and composed of thick-walled hyphae, but the inner layers are less dense, and have thin walls (Text-fig. 6). As the fertile hyphae grow and branch, the sterile ground tissue of the perithecium disintegrates from the centre

towards the periphery, until a few layers only of the wall remain. The cavity is occupied by the asci and by the remains of the ascogenous hyphae (Pl. XIII, fig. 6), the walls of the asci also disappearing as the fruit body reaches maturity. The ascospores lie free in the lumen of the ripe fruit, surrounded by a wall three to four layers thick. They are dispersed by the decay of the wall, or by its rupture.



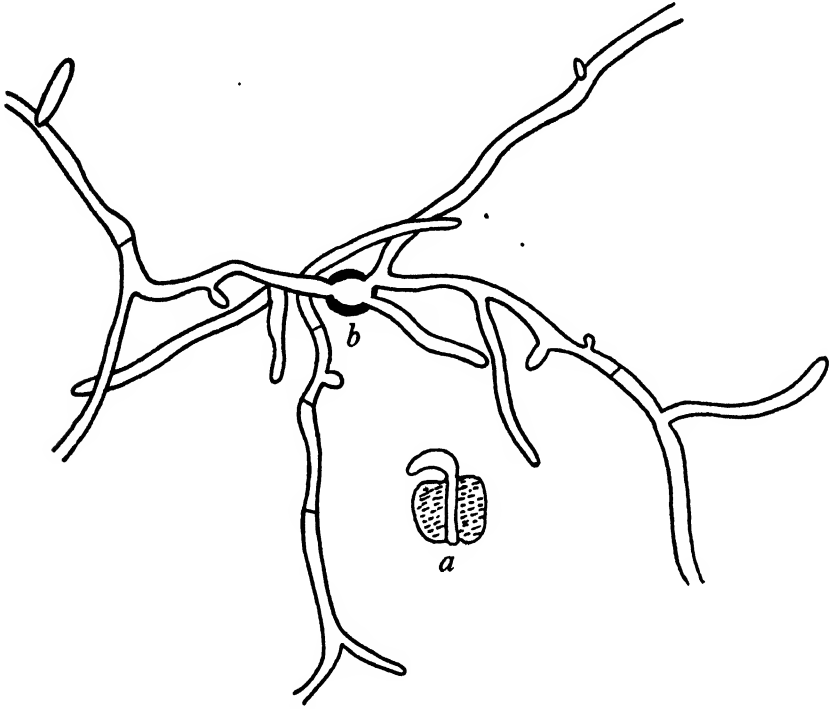
Text-fig. 6. Sclerotic perithecial wall and remains of ascogenous hyphae. $\times 1500$.

The ascospores are barrel-shaped, surrounded by a wall consisting of a thin inner layer and a thick outer layer. There is a longitudinal furrow in the outer wall, bordered by two more or less prominent ridges. At germination, the ascospore swells and puts out a germ tube opposite the furrow (Text-fig. 7).

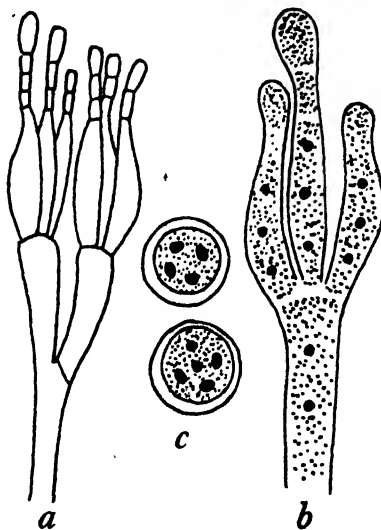
Perithecia may be obtained from cultures started from a single conidium or a single ascospore; *Penicillium egyptiacum* is homothallic.

Mycelium and conidiophores. The hyphae consist of multinucleate segments with strongly vacuolated contents, septa being usually

widely spaced. The segments of the conidiophores, as well as the metulae, are multi-nucleate (Text-fig. 8), and the conidia contain four or five nuclei (Text-fig. 8c). At germination, the conidium



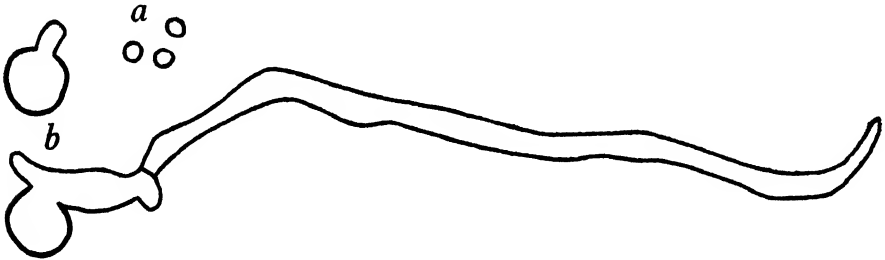
Text-fig. 7. Germinating ascospores. *a*, $\times 1500$; *b*, $\times 500$.



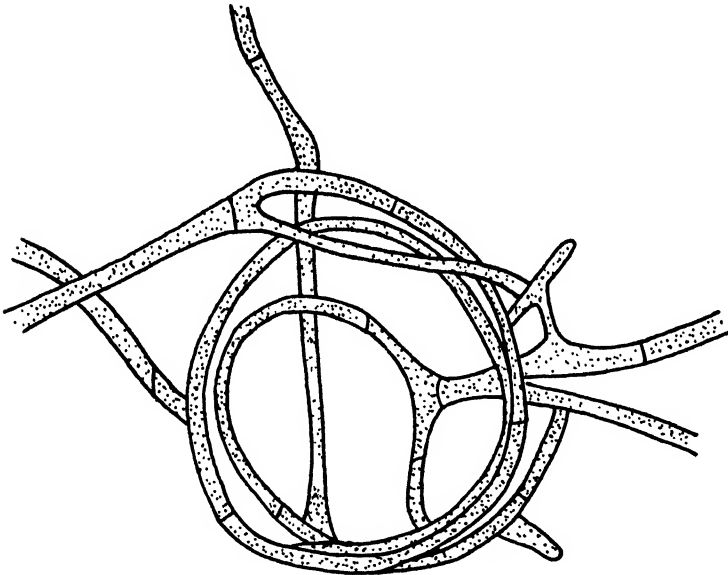
Text-fig. 8. Conidiophores and conidia. *a*, $\times 1500$; *b*, $\times 2500$; *c*, $\times 3000$.

swells to about three times its original size, and puts out one or more germ tubes (Text-fig. 9).

Occasionally flat coils of hyphae occur in the cultures (Text-fig. 10). They bear conidiophores, and they do not give origin to perithecia.



Text-fig. 9. *a*, conidia before germination; *b*, during germination. All $\times 1000$.



Text-fig. 10. Flat mycelial coils. $\times 2500$.

CONCLUSION

It is not necessary to utilize special conditions of culture to obtain the perithecia of *P. egyptiacum*. This species grows on a wide range of substrata. When these are of low nutritive value, the colonies are thin, and the conidiophores have a special tendency to form on rope-like strands of the aerial mycelium; perithecia develop in very small numbers on poor media. When there is a good supply of carbohydrates, vegetative growth is strong and perithecia abound. It seems likely that the conditions which favour the growth of the perfect stage depress conidial development, but the reverse is not true, for

perithecia were seen on every medium on which the fungus would grow. They began to appear, usually, during the second or third day after the culture was started, at 25° C. By the end of the week cultures were distinctly brown with perithecia.

P. egyptiacum differs considerably from the species described by Brefeld (3), Dangeard (7) and Fraser & Chambers (9), since it neither possesses an antheridium nor is the archicarp coiled. It closely resembles *P. Ehrlichii*, described by Klebahn (12), which has a perithecial rudiment consisting of a thick, nearly straight septate hypha, and has no antheridium.

The rapid formation of ascogenous hyphae by *P. egyptiacum* distinguishes the species clearly from the members of the *P. "crustaceum"* group, in which, according to Brefeld (3), asci develop in a sclerotium-like body only after a long period of rest. Klebahn (12) mentions that Morini (15) found that the sclerotia of *P. candidum* begin to contain asci about a fortnight after the development of the sclerotium, and that the ripe perithecia have a thin few-layered wall, covered externally by the granular remains of the disorganized outer envelope, and enclosing a cavity wholly filled by intact asci with ascospores. Klebahn noted that the perithecia of *P. Ehrlichii* mature rapidly and have a two-layered wall.

In *P. egyptiacum* the fruit body begins as a compact mass in which ascogenous hyphae may be found. They branch and develop without any period of rest, and soon give asci. The mature perithecial wall is three or four layers thick, and its cavity contains free ascospores. There is no envelope. In general, this species, apart from the absence of an inactive period, agrees with the "truffle" type of Sopp (20).

Gäumann (10) distinguishes two types of ascogenous hyphae in the genus. The first occurs in *P. vermiculatum*, where the ascogenous hyphae are converted into a chain of binucleate cells without any arrest in development, these yielding asci after a nuclear fusion. The second occurs in *P. crustaceum*. In that species inconspicuous aseptate ascogenous hyphae lie inside the hard sclerotium, but, some two months after the sclerotia are sown, the ascogenous hyphae become segmented into stout, cylindrical, and apparently binucleate segments. There is further complicated branching before the asci finally appear.

In *P. egyptiacum* the ascogenous hyphae are cut into a chain of binucleate segments by septa appearing at a late stage of development, but the nuclear behaviour has not yet been worked out.

Penicillium egyptiacum presents many points of interest, and further work is needed to complete many of the experiments briefly described in this preliminary account.

SUMMARY

1. *Penicillium egyptiacum* isolated from soil in Egypt, differs from most species of the genus in producing perithecia with readiness on most common media.

2. Tests of its behaviour at various temperatures, on media of different pH, in conditions of different humidity and at reduced atmospheric pressure, show that special conditions of the environment have no particular effect on the formation of the perithecia. Of the conditions tested, humidity seems to exert most effect.

3. The archicarp is not coiled, but has the form of a short, stout septate hypha.

4. Asci develop rapidly, and not after a period of inactivity.

5. The species is homothallic.

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EXPLANATION OF PLATE XIII

- Fig. 1. *Penicillium egyptiacum* grown on Barnes's medium.
- Fig. 2. *P. egyptiacum* grown on Richard's sol. agar.
- Fig. 3. Sectors on malt agar.
- Fig. 4. Saltant with very few perithecia and tremendous crop of conidia.
- Fig. 5. Young perithecium filled with sterile tissue: the remains of ascogenous hyphae are darkly stained near the centre. $\times 150$.
- Fig. 6. Perithecia in course of development. $\times 136$.



Fig. 1



Fig. 2



Fig. 3



Fig. 4

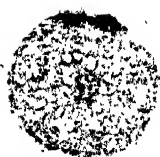


Fig. 5



Fig. 6

THE ORIGINAL AND MODERN CONCEPTIONS OF *STEMPHYLIUM*

By S. P. WILTSHIRE

Imperial Mycological Institute, Kew

(With Plates XIV and XV and 17 Text-figures)

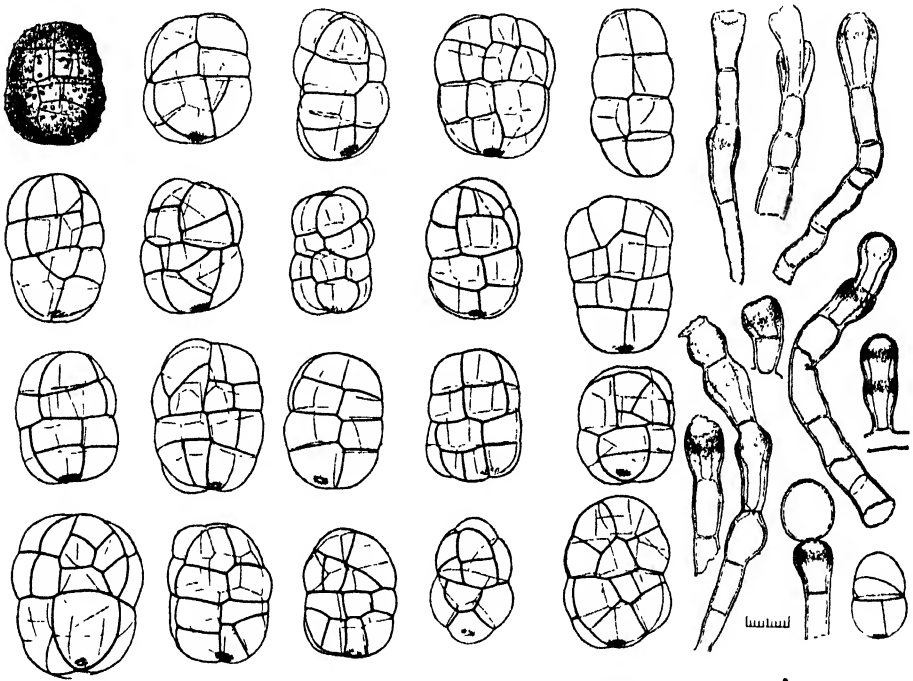
THE ORIGINAL CONCEPTION OF *STEMPHYLIUM*

THE genus *Stemphylium* was founded by Wallroth in 1833 on the single species *S. botryosum*. The description of the genus reads: "Hyphae [i.e. conidiophores] simplicissimae breves articulatae nodulosae, vertice incrassato sporidium ovatum subangulatum longitudinaliter transversimque septatum, veluti multiloculare sustinentes." This implies fungi with the conidiophore swollen at the apex and with "short nodular segments" bearing an ovate, subangular, muriform spore. The diagnosis of the species reads: "Hyphis articulato-nodulosis abbreviatis erectis in fasciculos distinctos gregatimque excurrentes junctis fragilibus, sporidio ovato subangulato longitudinaliter transversimque septato, veluti concamerato laxe appenso opaco nigro majusculo terminatis." The nodular segments of the conidiophore and the ovate, subangular shape of the muriform spore are again mentioned, and though the swollen apex of the conidiophore is not referred to, the description of the nodular segments clearly shows that such a swelling must have occurred, as the nodular segments are nothing more than the succession of swollen apices.

The specimen from which Wallroth very probably described the species is preserved in Herb. Wallroth at Strasbourg University, and through the kindness of Prof. Chermezon I was permitted to borrow it for examination. It consists of four pieces of stem gummed on a small card bearing the label "*Stemphylium botryosum* W. ad *Asparagum*" presumably in Wallroth's handwriting (Pl. XIV, fig. 1). The information given by the label is meagre, but Prof. Chermezon remarks that this is usual in Wallroth's herbarium. The host, asparagus, is the "type host" for the species, the original collection having been recorded on dried stems of asparagus.

The fungus agrees entirely with that indicated by the diagnosis (Text-fig. 1). The conidiophores are fasciculate, in groups ranging from three or four to upwards of fifty, apparently arising from an immersed stromatic cushion, or sometimes sparsely on a superficial mycelium. They are dark in colour, occasionally branched, septate

at intervals of about 9 or 10 μ , mostly from 30–70 or 80 μ long and 5–7 μ wide but swollen at the apex to a diameter of 7–10 μ . The apical swelling is darker below than in the upper half and is thickened by an inner wall. The growth of the conidiophore through the apex, giving rise to a succession of “nodular segments”, is clearly recognizable; up to four swellings were observed, the distance from the top of one to that of the next varying from 10 to 50 μ . The organization of the conidiophore corresponds with that described below for *Macrosporium sarcinaeforme* and already recorded by Miyake for *M. parasiticum*. The conidia are dark, verrucose, roughly oval or subangular,



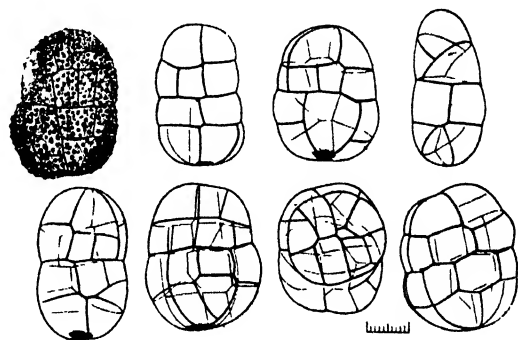
Text-fig. 1. *Stemphylium botryosum* Wallroth. Conidiophores and conidia drawn from Wallroth's specimen. $\times 500$.

slightly constricted at the median transverse septum and to a lesser degree at the other cross-walls, muriform with transverse, vertical and oblique septa dividing the conidium into many cells, with a prominent basal scar, and measure 24–39 \times 19–31 μ .

THE IDENTITY OF *STEMPHYLIUM BOTRYOSUM* WALLROTH WITH THE CONIDIAL STAGE OF *PLEOSPORA HERBARUM*

There can be little doubt that *Stemphylium botryosum* Wallroth is the fungus which has been known in the literature as *Macrosporium sarcinula* Berk. (or *M. parasiticum* Thüm.) the conidial stage of *Pleospora herbarum* Rabenh.

Berkeley described *Macrosporium sarcinula* in 1838 on decaying orange gourds, collected at King's Cliffe. The type material of *M. sarcinula* in Herb. Kew. consists of a small, rather curled up piece of orange rind, with one large blackish spot towards the edge, 9 mm. across in its maximum diameter; pinpricks indicate that another portion was formerly attached to the paper on which the specimen is mounted. It is labelled "*Macrosporium sarcinula*, Berk. Ann. of N.H. King's Cliffe, ex herb. Berk." in Berkeley's handwriting. There are five other specimens of Berkeley's in the folder, but those from King's Cliffe, the type locality, are of a later date than 1838. The two fungi confused by Berkeley in his original description of the fungus (Pl. XIV, fig. 2) are present on the specimen, viz. one "with short clavate filaments" (a species of *Alternaria*), and the other with spores which have "a rectangular outline and resemble very much little corded bales, from which circumstance the name is taken". Berkeley considered the former to be the immature stage of the latter, but this has not proved to be so. Only a very few sarcinaeform conidia were found in preparations made from the type, whereas large numbers of the *Alternaria* occurred. The sarcinaeform conidia obtained from the specimen (Text-fig. 2) were ovoid-oblong with a constriction at the median transverse wall, light brown, covered with small discrete warts and marked with a conspicuous basal scar; the measurements of the few spores seen ranged from $31-35 \times 20-26 \mu$. They agree very well with those of the type specimen of *Stemphylium botryosum*.



Text-fig. 2. *Macrosporium sarcinula* Berk. Conidia drawn from the type specimen. $\times 500$.

The conidia observed did not represent the full range of variation which they attain, for Berkeley figured them much longer, more angular, and much more resembling "bales of cotton" in shape (Pl. XIV, fig. 2). No conidiophores were observed in the type material, but Berkeley remarked that "a few of the peduncles [conidiophores] are seen amongst the sporidia, their articulations being swollen above", and his figure agrees with those of *S. botryosum*.

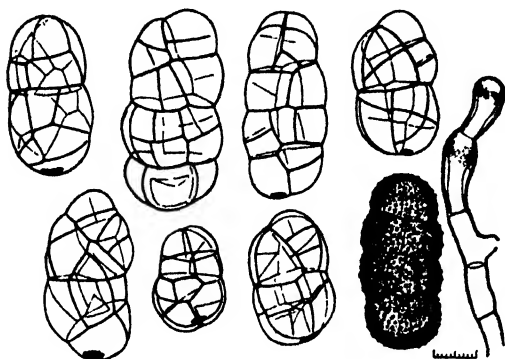
The association of *Macrosporium sarcinula* with *Pleospora herbarum* as its ascigerous stage really dates from the Tulasnes who, in 1863, cited the former (among various other species) as the conidial stage of the latter and also figured germinating ascospores bearing the

characteristic conidiophores with conidia attached. Gibelli & Griffini in 1872 named the ascigerous stage *P. sarcinulae* (= *P. herbarum*), and they also illustrated an ascospore bearing two conidiophores and conidia attached. Miyabe in 1889, studying the fungus on onion, proved the genetic connexion between the conidial stage, named *Macrosporium parasiticum* by von Thümen in 1877 (Myc. Univ. 667), with *Pleospora herbarum* by pure cultures and expressed the opinion that *Macrosporium sarcinula* Berk. was the same fungus, though he did not examine type material of this species. Prillieux & Delacroix (1893) doubted whether *M. parasiticum* could be maintained as a species distinct from *M. sarcinula*, and suggested it may form the conidial stage of *Pleospora herbarum*; they were evidently not aware of Miyabe's paper. Hanzawa (1915) suggested that the seven species of *Macrosporium* recorded by Saccardo on *Allium* (including *Macrosporium sarcinula*) are all perhaps identical with, or nearly related to *M. parasiticum*. Teodoro (1923) considered that Miyabe had established the identity of *M. parasiticum* with *M. sarcinula*, but throughout his paper he refers to the onion parasite as *M. parasiticum*, which he confirmed as the conidial stage of *Pleospora herbarum*. Kidd & Beaumont (1924) obtained the conidial stage (which they call *Macrosporium commune* Rabenh.) with its ascigerous stage, *Pleospora herbarum*, in culture. Bolle (1924) maintained that *Macrosporium parasiticum* and *M. sarcinaeforme* (which she regarded as identical with it) are merely strains of *Pleospora herbarum*, for the conidial stage of which the name *Macrosporium sarcinula* should be retained. Machacek (1929) & Ellis (1931) confirmed the relation between *M. parasiticum* and *Pleospora herbarum*. Verwoerd & Du Plessis (1931) stated that *Macrosporium sarcinula* is the conidial stage of *Pleospora herbarum* on the onion.

Von Thümen's exsiccatum of *Macrosporium parasiticum* comprises excellent material from which Text-fig. 3 has been drawn. The conidia are longer than observed in the type of *M. sarcinula*, but, taking into account Berkeley's figure, there is very strong evidence that the two species are identical. The spores of *M. parasiticum* are also longer than those of *Stemphylium botryosum*, but nevertheless I would regard both as belonging to the same species and, consequently, *Macrosporium sarcinula* and *M. parasiticum* as synonyms of *Stemphylium botryosum*. The fungus is common, occurring on a wide range of hosts and very probably has been known under a number of names. In culture the conidia are definitely smaller than on the host as recorded by Bolle (cf. Bolle, 1924, p. 63 and Pl. III, figs. F, G, H, I). The extreme measurements of conidia collected on dying onion leaves at Oakley, Hants, in 1932 were $21-49 \times 10-22 \mu$, whereas in culture they were $20-26 \times 12-18 \mu$, the measurements indicating a change of shape as well as of size. Seeing such variation occurs in culture it may also be expected to occur on different hosts in nature, and minor differences

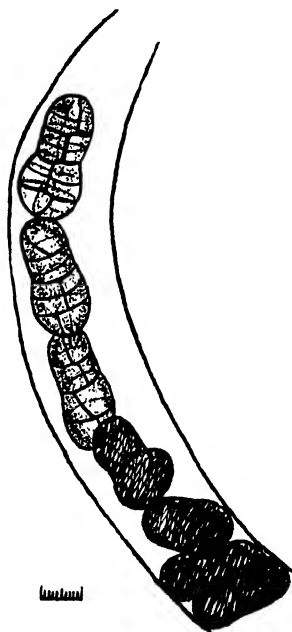
in spore size alone cannot therefore be regarded as satisfactory grounds for maintaining species.

Confirmatory evidence of the correct identification of the conidial stage is frequently afforded by the development of the ascigerous stage (*Pleospora herbarum*) with its characteristic ascospores with seven cross-septa (cf. Mason, 1928). Isolations of the fungus from various hosts (including lettuce, tomato, flax, sisal, olive, orange, lemon) have been found almost always to form perithecia if the cultures are kept in the light. The importance of light was demonstrated by one Petri dish culture, of which a portion shaded by a label was observed



Text-fig. 3. *Macrosporium parasiticum* Thüm. Conidia and conidiophore from Thümen's material (Mycotheca univ. 667: on leaves of *Allium cepa*). $\times 500$.

Text-fig. 4. Immature ascospores of *Pleospora herbarum* Rabenh. found in perithecia associated with *Stemphylium botryosum* Wallr. on the type specimen of the latter species. $\times 500$.



Text-fig. 4.

not to bear perithecia, whereas these occurred plentifully in the unshaded parts. Some strains tend to develop the perithecial stage more freely than the conidial, whilst others do just the reverse, and others again form both equally readily, but the comparative ease with which both stages may be obtained in culture will probably assist very greatly in elucidating the synonymy of each.

Similar evidence of the identity of *Stemphylium botryosum* as the conidial stage of *Pleospora herbarum* is afforded by the type specimen itself. In his remarks on *Stemphylium botryosum* Wallroth mentioned the association of *Sphaeria complanata* Tode with it, and this probably refers to the scattered perithecia which are conspicuous on the type. These are similar in appearance to those of *Pleospora herbarum*. They are mostly very immature, but some contained asci (Text-fig. 4) with

dark-coloured spores, mostly divided by three transverse and some longitudinal septa, but in one ascus seven cross-septa were recognizable in two of the ascospores (though one spore had the seventh septum rather indefinite), and in the other ascospores the number of cross-septa was not ascertainable though clearly more than three. The mature spores were evidently muriform, with seven cross-septa. Thus the search for the ascigerous stage in the type material was rewarded by finding what I believe to be the immature stage of *P. herbarum*, confirming the conclusion arrived at from the conidia alone.

The synonyms of *Stemphylium botryosum*, if my conception of the specific limits is correct, will probably be comparatively numerous. No attempt has been made to list these, but the following are given, as type material of each has been examined. Those occurring on onion are evidently synonymous with *Macrosporium parasiticum* and consequently, I believe, with *Stemphylium botryosum*.

Septosporium scyphophorum Cooke & Harkness in *Grevillea*, ix, 129, (1881). [Type on *Eucalyptus globulus* in Herb. Kew. Text-fig. 5.]

Mystrosporium alliorum Berk. apud Saccardo, *Syll.* iv, 541 (1886). [Type on onion in Herb. Kew. The citation of this species in the *Sylloge*, *Gdnrs' Chron.* p. 192 (1878), only refers to an answer to a correspondent by M. J. Berkeley in which he says he found on onions an undescribed fungus belonging to the genus *Mystrosporium*. The species was also mentioned as occurring on onions at Culver, Exeter, by Berkeley and Broome in *Ann. Mag. nat. Hist.* Ser. 5, ix, 183, No. 1982 (1882), but without diagnosis. Saccardo's description is the first published and the species must date from then, Berkeley's name being a *nomen nudum*. Text-fig. 6.]

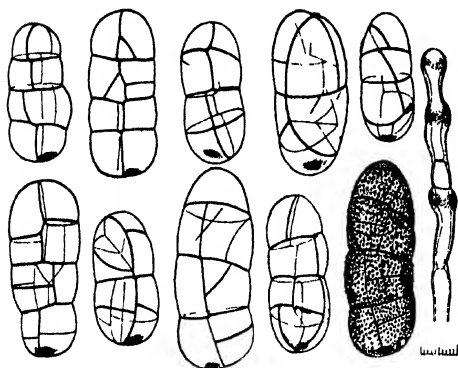
Macrosporium alliorum Cooke & Massee in *Grevillea*, xvi, 79, p. 80 (1888). [Type on onion in Herb. Kew. Text-fig. 7.]

Macrosporium Symplocarpi H. & P. Sydow in *Ann. Mycol., Berl.*, xi, 65, Fig. 5 (1913). [Type on *Symplocarpus foetidus* (Fungi exot. exs. 100). Specimen in Herb. Kew. examined. This species is very like *Stemphylium botryosum* Wallr. The spores are verrucose (sometimes indistinctly), light-coloured, and the cross-walls rather more prominently marked than usual. The conidiophores are light-coloured, with a wider swelling at the apex (10μ instead of about 8μ), and the thickening of the wall below the apex is very marked. These differences are not regarded as of specific rank and for the present the species is placed in *S. botryosum* Wallr. Text-fig. 8.]

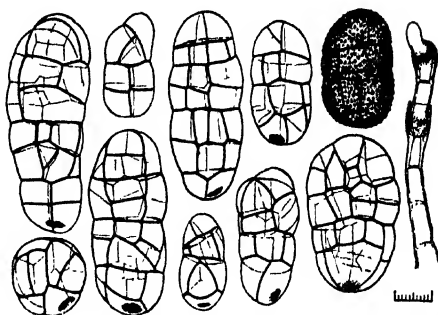
(?) *Thyrodochium Dracaenae* Werdermann in *Ann. Mycol., Berl.*, xxii, 189, 1 fig. (1924). [Type on *Dracaena draco* was kindly loaned to me by Dr Sydow. The species is discussed below, p. 223.]

Mason (1928) has already shown that Penzig's figure of *Macrosporium commune* Rabenh. refers to the wrong fungus. Cooke's original material of *M. Cheiranthi* Fr.β. *Betae* Cooke (Fungi Brit. Exs. 197 =

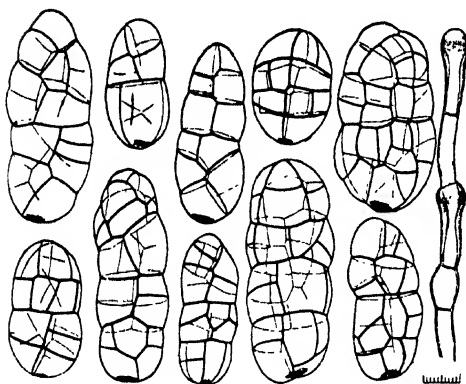
M. commune Rabenh., *Fungi Europaei*, 1360) contains an admixture of conidia of *Pleospora herbarum*, and it was conidia of this type that Penzig drew. Many identifications of *Macrosporium commune* therefore really refer to *Stemphylium botryosum*, e.g. Brefeld (1891) and Kidd & Beaumont (1924) as cited above.



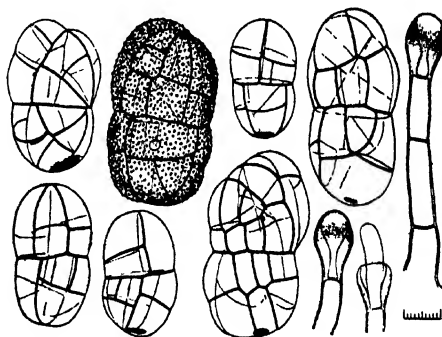
Text-fig. 5. *Septosporium scyphophorum* Cooke & Harkness. Conidia and conidiophore drawn from the type specimen on *Eucalyptus* bark in Herb. Kew. $\times 500$.



Text-fig. 7. *Macrosporium alliorum* Cooke & Massee. Conidia and conidiophore from the type specimen on onion. $\times 500$.



Text-fig. 6. *Mystrosporium alliorum* Berk. apud Sacc. Conidia and conidiophore from the type specimen on onion. $\times 500$.



Text-fig. 8. *Macrosporium Symplocarpi* H. & P. Sydow. Conidia and conidiophores from authentic material (*Fungi exot. exs.* 100) on *Symplocarpus foetidus*. $\times 500$.

THE CHARACTERS OF THE GENUS *STEMPHYLIUM* WALLROTH

The genus *Stemphylium*, as defined by its diagnosis, indicated by the diagnosis of its type species, and substantiated by its type specimen, is undoubtedly that named *Thyrospora* by Tehon & Daniels in 1924 and which in my paper on the foundation species of *Alternaria* and *Macrosporium* (1933) I referred to as group (c) comprising mostly species attributed to *Macrosporium*.

The characters which would suffice to distinguish the genus from other Phaeodictyae are (1) that the conidiophores are swollen at the apex which bears a single terminal spore (though this may be forced into a lateral position by the continued growth of the conidiophore); (2) that the growth of the conidiophore is continued through the terminal scar, the successive swellings recognizable in an old conidiophore marking the places where conidia have been borne; and (3) that the spore shape is oval or subangular, frequently constricted at the major, median transverse wall and never beaked. These characters are so clearly indicated in the diagnosis of the genus and its type species that I was able to recognize the genus implied before I knew of the existence of the Wallroth specimen.

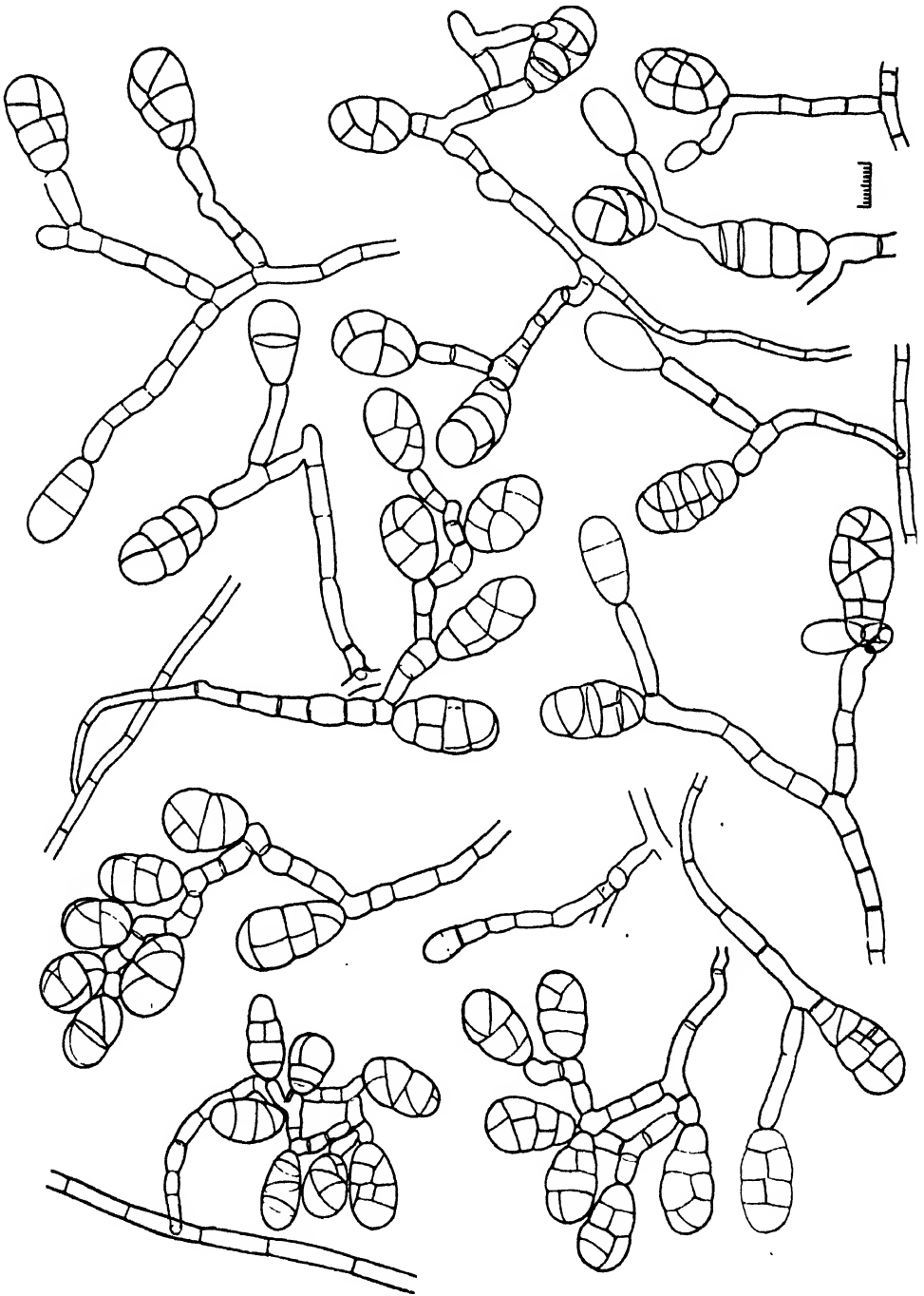
THE MODERN CONCEPTION OF *STEMPHYLIUM*

Elliott in 1917 expressed the opinion that species of *Macrosporium* with sarcinaeform or globose spores should be put into *Stemphylium* as "there is nothing in the morphology of the species of this group which would exclude them from the genus". He does not enlarge on the matter, and his views have not been generally accepted, especially as *Thyrospora* was erected for the species of *Macrosporium* in question. Elliott apparently was ready to widen the conception of *Stemphylium* to include the sarcinaeform *Macrosporium* species; he did not realize that these species must be placed in *Stemphylium*, as the genus was actually founded on a fungus of this type.

Bolle (1924) based her identification of *S. botryosum* on a culture isolated and named by Oudemans, which she regarded as agreeing with Wallroth's description. In order that the reader may understand the fungus Oudemans isolated, a figure of it, drawn from an authentic culture kindly supplied by the Centraalbureau voor Schimmelfcultures, is appended (Text-fig. 9).

Oudemans's fungus has no markedly swollen tips to the conidiophores, and the only resemblance it bears to the true *S. botryosum* is the shape of the conidia, but even these show no constriction at the median transverse septum. It is quite clear that Oudemans made a mistake in identifying his fungus with *S. botryosum*. Bolle interpreted *Stemphylium* in the sense of Oudemans and was convinced that it was distinct from *Macrosporium* which she would reserve for the species of the genus with sarcinaeform spores (viz. those for which Tehon and Daniels made the genus *Thyrospora*).

The present-day ideas of *Stemphylium* are generally those of Oudemans; the conidia are ovate and are borne either singly or in clusters on conidiophores which arise as branches from the mycelium. The first spore is terminal, and the conidiophore may then grow out laterally from the tip, not directly through the scar, to produce a



Text-fig. 9. *Stemphylium botryosum* sensu Oudemans. Conidiophores and conidia from cultures of Oudemans's strain. $\times 500$.

second spore, the first being pushed into a lateral position. This process is continued until a number of conidia are produced. The scars left by fallen conidia are recognizable on the sides of the conidiophore and exactly resemble those of *Alternaria*. Sometimes the conidiophore, instead of growing out immediately below the conidium, does so lower down so that it is branched. There is no marked swelling of the tip and no thickening of the lateral walls.

Seeing that the current views of the genus differ from the original, it may be interesting to trace the development of this change.

THE HISTORICAL DEVELOPMENT OF THE CURRENT VIEWS OF THE GENUS

No notice appears to have been taken of Wallroth's genus after it was erected in 1833 until Bonorden transferred eight additional species to it in 1851. Of these the first *Stemphylium pyriforme* was originally described by Corda as *Sporidesmium pyriforme*, and his figure is reproduced on Pl. XIV, fig. 3. Corda's fungus evidently had swollen tips to the conidiophores and bore more or less ovate conidia, one of which is figured, with a marked constriction at the median, transverse wall. It is a true *Stemphylium*, though from the description and figures it cannot be identified more precisely. Bonorden (Pl. XV, fig. 10) represented the fungus with pyriform spores and only slightly swollen tips to the conidiophores.

The second species *S. polymorphum* (= *Sporidesmium polymorphum* Cda) is represented by Corda (Pl. XIV, fig. 4) without swellings to the conidiophores and with irregularly shaped spores. Bonorden (Pl. XV, fig. 11) drew it as a true *Stemphylium*, but his figure does not agree with Corda's.

The remaining six species, in the absence of authentic material, appear to be of doubtful identity. None of them belongs to Wallroth's genus, judging by the original figures and diagnoses. *S. elegans* (Pl. XIV, fig. 5) (= *Sporidesmium elegans* Cda) may possibly be congeneric with Oudemans's *S. botryosum*, and *S. rhizospermum* (Pl. XIV, fig. 6) (= *Trichaeum rhizospermum* Cda) suggests an *Epicoccum*. *S. graminis* (Pl. XIV, fig. 7) (= *Soredospora graminis* Cda) is evidently a mixture of two fungi, the dictyosporous one having spores of irregular shape. The species has been recorded from apples and from oats, but both these records referred to fungi congeneric with Oudemans's strain of *S. botryosum*. *S. dubium* (Pl. XIV, fig. 8) (= *Mystrosporium dubium* Cda) is illustrated by a rather fanciful figure, and *S. bulbotrichum* (Pl. XV, fig. 9) (= *Septosporium bulbotrichum* Cda) with spores attached by stalks to swollen bases of the conidiophores. The last of the species mentioned by Bonorden, *S. macropodium* Cda, I have been unable to trace.

The fact that Bonorden transferred such a miscellaneous batch of species to *Stemphylium* testifies that he did not rightly appreciate the genus, and this conclusion is borne out by the species *S. sphaeropodium* which was described by him in 1864. Lindau (1908, p. 220) regards this species as of doubtful identity. The description accompanied the exsiccatum, Rabenhorst's *Fungi europaei* No. 689 which was collected by Bonorden on leaves of *Arundo* [= *Phragmites*] and comparison of this from Herb. Kew. with *Napicladium arundinaceum* (Corda) Sacc. (= *Helmi[ntho]sporium arundinaceum* Corda), as represented by Briosi and Cavara: I funghi parass. 419, indicated (as Mr Mason kindly suggested to me it would) that the species are identical. The conidiophore has a swollen base but not a swollen apex while the spores (which were not present on the specimen) are pyriform with two cross-walls but no longitudinal ones. Other synonyms of this species are as follows:

[*Hadrotrichum Phragmitis* Fuck., sensu Saccardo in *Mich.* II, p. 362 (1881), teste von Höhnelt, *Zbl. Bakt.* Abt. 2, LX, p. 8 (1923).]

Scolecotrichum Roumegueri Cav. 1890 in Briosi & Cavara; I funghi parass. 112 on *Phragmites communis*. [= *N. arundinaceum* (Corda) Sacc. teste von Höhnelt, *loc. cit.*] Exsiccatum in Herb. I.M.I., examined.

Napicladium laxum Bubák, 1906, in *Ann. Mycol., Berl.*, IV, p. 121 (1906) on *Phragmites communis*; type Kabat & Bubák, *Fungi Imperf. exs.* No. 48 [= *S. Roumegueri* Cav. teste von Höhnelt, *loc. cit.*].

Brachysporium Phragmitis Miyake, 1912, in *Bot. Mag., Tokyo*, XXVI, 303, p. 63, Figs. 10, 11 on *Phragmites communis*. [= *N. arundinaceum* (Corda) Sacc. testibus Shirai & Hara in *A List of Japanese Fungi*, p. 232 (1927).]

The next species to be placed in *Stemphylium* was *S. fuscescens*, a description and specimens of which were issued by Rabenhorst in 1866 in his *Fungi europaei*, II, No. 1174d. This exsiccatum contains masses of spores of *Pestalozzia* and a number of an *Epicoccum*. Ferraris in *Flora Italica*, p. 489 (1910) reduced the species to a variety (var. *fuscescens*) of *Stemphylium macrosporoideum*, but I was unable to find any spores resembling this species on Rabenhorst's specimens. Ferraris distinguished the variety on the ground of the warted spores but type material of *S. macrosporoideum* in Herb. Kew. has warted spores so that in any case the variety cannot stand. Furthermore, as shown later (p. 229), *S. macrosporoideum* is not a *Stemphylium* either in the sense of Wallroth or Oudemans.

In 1869 Corda's species *Sporidesmium paradoxum* was added to the genus as *Stemphylium paradoxum* by Fuckel, who issued an exsiccatum of the fungus on birch in *Fungi Rhenani* 1515. Corda's figure is very characteristic (Pl. XV, fig. 12). One of Fuckel's specimens preserved in Herb. Kew. shows a fungus with large multiseptate spores, rather irregular in outline with a smooth surface, fuscous except the end cell

which may be almost hyaline. It was difficult to determine how the spores were borne, but one showed (somewhat indistinctly) two hyphae attached to the base. These appeared to be ordinary hyphae and not specially developed conidiophores, and it is just possible that the spores may be bulbils though more material of the fungus is required for its structure to be properly investigated.

We now come to the important species *Stemphylium lanuginosum* which was described exceptionally fully by Harz in 1871, who also published a figure of the species (Pl. XV, fig. 13). This species is evidently the one upon which the current ideas of the genus are based. Harz obtained the fungus from honeycomb of a beehive affected by foulbrood, which had been placed for observation in a moist chamber. He remarks that fairly long lateral branches develop below a conidium, which again form a conidium at their apex, the same process being repeated below these branches so that a kind of cymose conidial system is formed. The conidia are stated to be unicellular at first but rapidly becoming divided by a transverse septum into two fairly equal halves; gradually turning brown it forms several transverse and longitudinal septa. The mature conidium is barely angular, oval, or rounded ovoid at both ends, light to very dark brown, 30μ in length and 21.5μ in breadth. This description agrees fairly well with Oudemans's *S. botryosum* ($9-39 \times 9-22\mu$ according to Bolle) but the conidia are slightly less elongated. There are no apical or nodular swellings to the conidiophores and also no median constriction in the spores. The characteristic branching of the conidiophores described by Harz occurs in Oudemans's strain whereas it has not been noticed in a number of cultures of allied species. Tentatively therefore I am inclined to consider *S. botryosum* of Oudemans as identical with *S. lanuginosum*. Saccardo renamed "*Mystrosporium lanuginosum* Harz" *Alternaria lanuginosa* in Syll. iv, 546 (1886), but as explained in Syll. xviii, 624 (1906), the former name was a *lapsus calami* for *Mystrosporium hispidum* which should be called *Alternaria hispida* (Harz) Oud.; no such species as *Mystrosporium lanuginosum* was described.

Although the fashion set by Harz has been followed by most systematists, no doubt partly by reason of the excellent figure Harz published, there are instances where the original usage has been adhered to. Horne & Horne (1920) described the conidial stage of *Pleospora pomorum* (= *P. herbarum*) as "of the *S. pyriforme* type", and Weber, in doubt about the correct genus for his *Stemphylium Solani*, decided to follow Elliott, and his species, erected in 1930, is a good *Stemphylium* in Wallroth's sense.

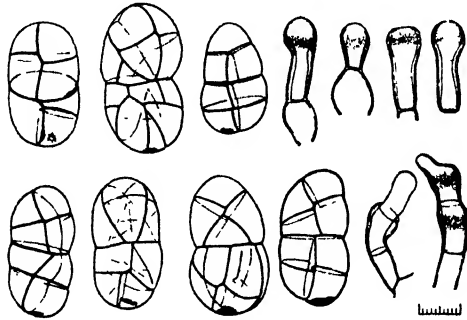
MODIFICATION OF THE ORIGINAL CONCEPTION OF *STEMPHYLIUM*
TO INCLUDE THE CURRENT ONE

Under the International Rules of Botanical Nomenclature, there seems to be no alternative to retaining Wallroth's characters for the genus *Stemphylium*. To avoid the necessity of transferring the species of *Stemphylium* sensu Harz to another genus I propose that both groups should still be included in the genus (the essential character of which would be the dark-coloured, muriform, ovate or subangular spore) and that they should be termed *Eustemphylium* and *Pseudostemphylium*, respectively, the former including those species with the original characters of the genus and the latter those allied to *Stemphylium lanuginosum* Harz. Ultimately it may be necessary to transfer the *Pseudostemphylium* section to a new genus, but to do so now would probably lead to great confusion as the genus is chiefly composed of members of this group. It was hoped that *Mystrosporium* might be the proper genus for the *Pseudostemphylium* group, as Bolle regarded it as synonymous with *Stemphylium*. It was founded on *Mystrosporium dubium* (Pl. XIV, fig. 8), and neither the generic diagnosis nor the description of the single type species affords any idea of the fungus concerned. The vague characters of the genus are sufficiently manifested by the fact that it has been regarded as synonymous with *Alternaria* by Elliott, with *Stemphylium* by Bolle, and with *Macrosporium* by Shear & Clements.

It may be remarked here that De Sousa da Camara in 1930 proposed the separation of the genus *Soreymatosporium* from *Stemphylium* on the basis of its pleurogenous conidia, the latter genus being retained for those species with conidia of the acrogenous type. In both *Eustemphylium* and *Pseudostemphylium* groups the spores arise terminally and may be pushed into a lateral position if the conidiophore continues growth. In his description of *Stemphylium dendriticum*, Da Camara gives the spores as acro-pleurogenous, but his Figs. 4, 5 and 6 suggest the usual method of formation, with which the drawing in his Fig. 7 does not seem to correspond. Pleurogenous conidial formation appears to be very rare in species of *Stemphylium*, and its occurrence indeed requires confirmation.

Another genus which must be considered in connexion with *Stemphylium* is *Thyrodochium* founded by Werdermann on *T. Dracaenae* in 1924, type material of which was kindly lent me by Dr Sydow. This species has conidia and conidiophores exactly similar to *Stemphylium* Wallr., but the conidiophores are borne in a cluster on a stroma which induced Werdermann to place it in the Tuberculariaceae. But *S. botryosum* itself also occurs sometimes on a stroma, and in fact the type material of this species showed equally marked stromatic development as the type of *Thyrodochium Dracaenae*. In shape and

size the spores (Text-fig. 10) are very like *Stemphylium botryosum* Wallr. and I think that *Thyrodochium Dracaenae* is really synonymous with it. *Thyrodochium* antedates *Thyrospora* by a year, so that in any case it is extremely doubtful whether the latter genus could stand, and by interpreting *Stemphylium* as I have suggested we have a very old and presumably stable genus in which to place these species.



Text-fig. 10. *Thyrodochium Dracaenae* Werderm. Conidia and conidiophores drawn from type material on *Dracaena draco*. $\times 500$.

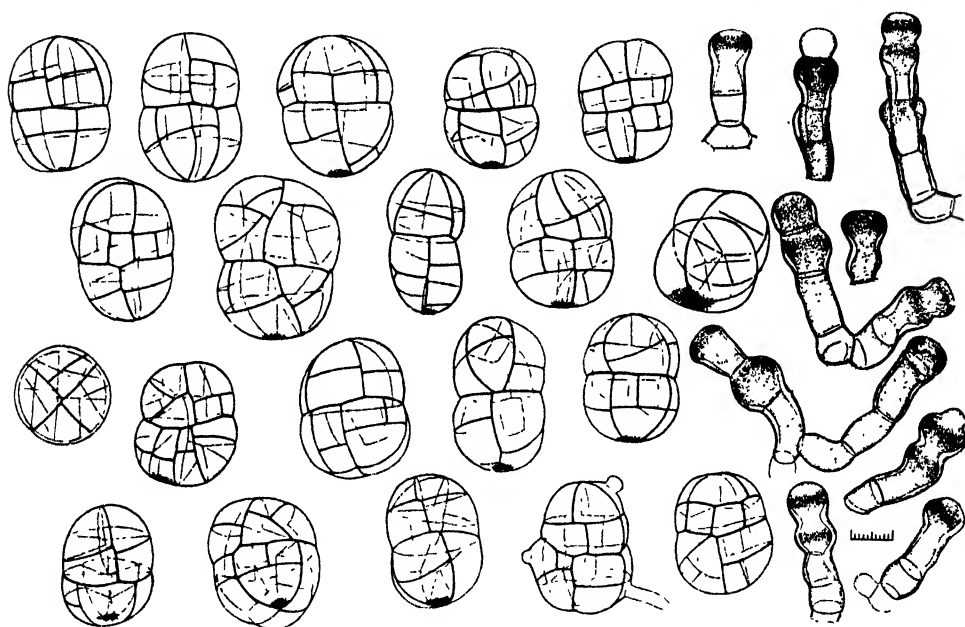
TRANSFERENCE OF SPECIES OF *THYROSPORA* TO *STEMPHYLIUM*

Thyrospora sarcinaeforme (Cav.) Tehon & Daniels

To *Eustemphylium* must be transferred the species now placed in *Thyrospora*. Of these the first is *T. sarcinaeforme*. As Bongini has recently shown, Tehon & Daniels in erecting *Thyrospora* on *Macrosporium sarcinaeforme* Cav. did not correctly identify their fungus, for it had echinulate spores whereas *M. sarcinaeforme* has smooth spores. I collected Cavara's fungus on clover at Oakley, Hants, in September, 1932, where it occurred extensively in a field, producing on the leaves dark brown, zonate spots, ranging from 1 to 8 mm. in diameter, broadly oval or circular (Pl. XV, fig. 14). The conidiophores (Text-fig. 11) occurred singly for the most part but were sometimes grouped; they projected more or less perpendicularly from the epidermis and measured from 16–50 μ long by 6–8 μ in diameter at the base and 9–12 μ at the apical swelling. Close examination of the conidiophore showed that sometimes the apical swelling or nodular segments arose from within the lower part of the hypha, through the dark outer wall of which the inner cell wall could be recognized.

Method of conidial formation. As the method of formation in *Stemphylium* is an important character detailed observations of the process in this species were made. It was found to correspond with that described by Miyabe (1889, p. 2) for *Macrosporium parasiticum*. In culture the actively growing, vegetative hypha, when about to form a conidiophore (Text-fig. 12), swells up towards the end which

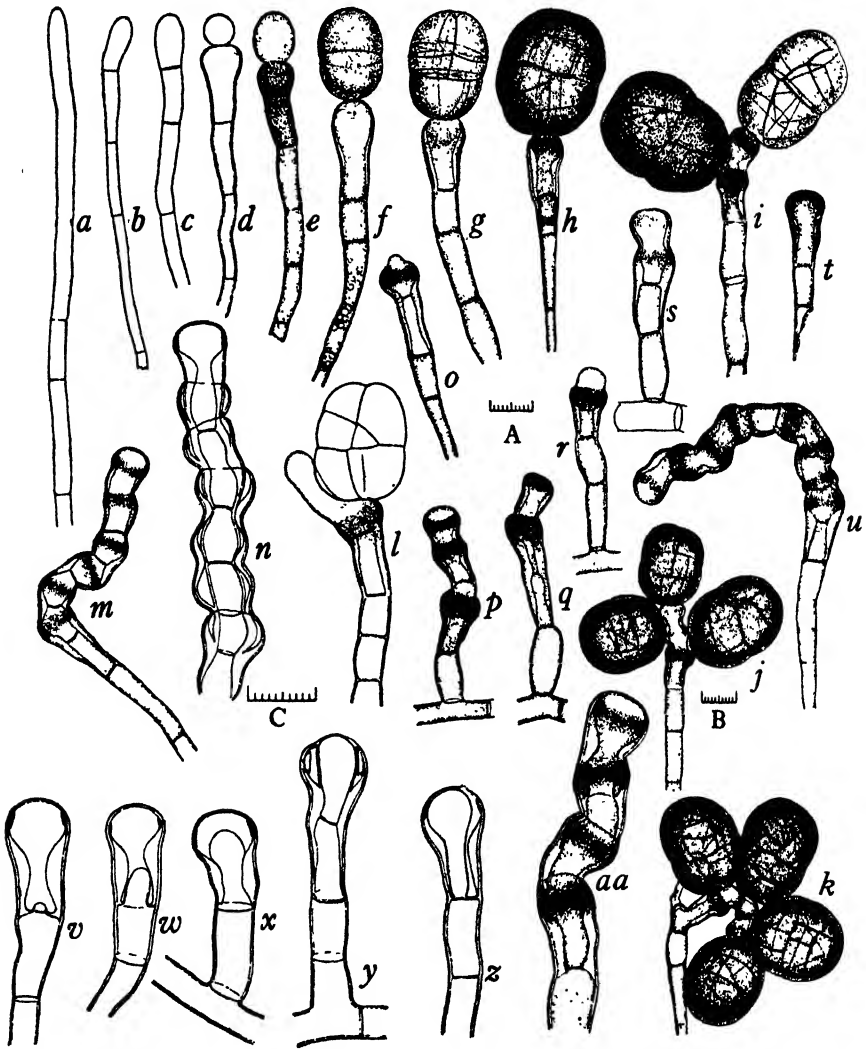
becomes somewhat club-shaped. A small, thin-walled, spherical or slightly oval, hyaline body, the young spore, is formed on its apex, which is often rather flattened, and the sides of the terminal cell of the conidiophore become slightly thickened. The immature spore increases in size, but the wall remains thin whilst at the same time the thickening of conidiophoral wall becomes more apparent. This thickening gradually increases until a well-defined greenish or hyaline inner layer is formed, chiefly in the lower part of the terminal cell and reaching its greatest thickness just below the swelling, above which it rapidly thins again so that there is no thickening visible at the apex.



Text-fig. 11. *Stemphylium sarcinaeforme* (= *Macrosporium sarcinaeforme* Cav.). Conidia and conidiophores from clover; the top three rows from material collected at Oakley, Hants, in September 1932, the bottom row from Cavara's exsiccatum, I funghi exsicc. No. 116. $\times 500$.

When rather more than half grown the spore divides into two approximately equal cells by a transverse wall, a slight constriction being noticeable at this wall at a very early stage. Vertical, cross, and oblique walls are then formed and the spore deepens in colour as does also the conidiophore, especially at its swollen part. The mature spore is a heavy structure for the conidiophore to bear and the thickening of the sides of the latter may be on account of this. Just below the region where the spore is attached is a somewhat broad, darkened zone, suggesting in optical section a bubble of gas, whilst at the base of the conidium itself a dark "bubble" is often seen. When the spore is mature it may fall off, or the conidiophore continue

development with it still attached, forcing the spore into a lateral position. There are two methods by which the conidiophore may continue growth (*a*) simply by the growing out of the apex through



Text-fig. 12. *Strophylium sarcinaeforme* (= *Macrosporium sarcinaeforme* Cav.) in culture, showing the development of the conidiophores and conidia. *a-h*, early stages in the formation of conidiophores and conidia. *i-k*, formation of two, three, and four conidia on single conidiophores. *l*, conidial formation within the agar and continued growth of the conidiophore. *m-u*, different types of conidiophores. *v-aa*, stages of growth of intrahyphal hyphae on the septal wall of the terminal cell of the conidiophore. Magnification, all $\times 500$ (scale A); except *j*, *k*, *m*, *o-u*, \times approx. 400 (scale B); *n*, *v-aa*, $\times 800$ (scale C).

the scar, or (*b*) by a new hypha growing up from the septal wall below, through the apical swelling (where it sometimes becomes swollen), and piercing through the apical scar forms a new apical

swelling. Such a hypha is thin-walled and difficult to discern when young, but with age it gradually thickens and it can be readily recognized in old conidiophores. The transverse wall from which the growth takes place may be the original basal wall of the terminal cell or a new transverse wall, formed usually where the inner thickening is greatest. These two methods are often seen in one conidiophore; with the first there is no fracture of the conidiophoral wall such as can be discerned, though sometimes with difficulty, with the second.

An explanation of the existence of the two methods may be found in the wounding of the conidiophore. Buller (1933, pp. 130-40) has shown that when a hyphal cell is wounded, the perforation in each septal wall bounding it is immediately plugged and within a short time a new hypha grows out from the plugged wall. It is quite conceivable that the spore, either in falling off or as the result of movement, may injure the conidiophore, and when this occurs a fresh hypha grows up from the septal wall below to continue the development of the conidiophore. Where no injury takes place then the growth continues from the apex uninterrupted. It may be objected that if such an explanation is correct the two methods of conidiophore growth should occur in other fungi. As a matter of fact, apart from *Macrosporium parasiticum* they have been recorded already by Wardlaw in *Helminthosporium torulosum*, whilst I have also seen them in various species of *Alternaria* where, however, the intrahyphal hyphae are difficult to make out as there is no apical swelling to the conidiophore. Duvernoy and Maire have also described a modified intrahyphal growth of the conidiophore in *Endophragmia mirabilis*. It would not be surprising if both methods occur in a number of big-spored fungi.

The conidiophore of *Macrosporium sarcinaeforme* may continue to develop until a number of spores are formed in a cluster. The positions of the spores can often be recognized by the flattened bends in the conidiophore, but sometimes the apical swelling appears to grow out into another apical swelling without bearing a spore.

The mature conidia are ovate but with a median constriction, each half approximating to rather more than half a sphere. Sometimes the conidia are rather elongated, reaching to nearly twice as long as broad; they are much divided, olivaceous in colour but remain somewhat translucent and have a smooth surface. The point of attachment is marked by a basal scar which is very well developed and consists of a darkening of the cell wall round a small light-coloured area. In size the conidia measure $28-38 \times 18-29 \mu$ on the host-plant and $21-36 \times 16-28 \mu$ in culture. Cavara's measurements were $24-28 \times 12-18 \mu$, but evidently he measured smallish, possibly immature conidia (as may be gathered from his figure), for on his *exsiccatum* (I funghi parass. No. 116) they were $28-31 \times 21-25 \mu$ (Text-fig. 11). Though there is a slight divergence in spore measure-

ments between the Oakley collection and Cavara's material the general similarity of the fungi is so close as to leave no doubt of their specific identity. Through the kindness of Prof. McCallan a culture of his isolation of *M. sarcinaeforme* from red clover was received from the United States, and this agreed with the Oakley fungus. Bolle (1924, p. 68) considered that *M. sarcinaeforme* was identical with *M. parasiticum*, but the strains she regarded as *M. sarcinaeforme* were isolated from seeds of *Trifolium* and *Medicago* supplied by Dr Gentner who in 1918 stated that *Macrosporium sarcinaeforme* was the conidial stage of *Pleospora herbarum*. Both Gentner and Bolle appear to have misidentified *Macrosporium sarcinaeforme* and to have really had a strain of *Pleospora herbarum*.

The identity of the species *Macrosporium sarcinaeforme* Cavara, therefore, is clear; it evidently belongs to the *Eustemphylium* section of the genus *Stemphylium* and is accordingly renamed *S. sarcinaeforme* (Cav.) comb.nov., and the name *Thyrospora sarcinaeforme* (Cav.) Tehon & Daniels exd.spec. then becomes a synonym.

Turconi & Maffei (1912) erected *Macrosporium Sophorae* for a fungus causing a spotting of leaves of *Sophora japonica*. An exsiccatum of this species was issued by Pollacci in *Fungi Longobardiae*, exs. 298, and this has proved to be *Stemphylium sarcinaeforme*. The authors state that their species differs from the latter in the shape and colour of the spots and the shape and dimensions of the conidiophores and conidia. The spots may be expected to differ somewhat from those of clover but the shape and dimensions of the conidiophores and conidia agree very well with the revised diameters for *S. sarcinaeforme* given above.

Thyrospora parasitica (Thüm.) Angell

The second species to be placed in *Thyrospora* was *T. parasitica* (Thüm.) Angell, 1929. This species was originally named *Macrosporium parasiticum* Thüm. which has already been shown to be synonymous with *Stemphylium botryosum* Wallroth.

Thyrospora Solani (Weber) Sawada

The third species to be placed in *Thyrospora* was *T. Solani* described by Weber in 1930 as *Stemphylium Solani* and transferred to *Thyrospora* by Sawada in 1931. It is a true *Thyrospora* but as this genus is here treated the original name is the valid one. The species is said to have spores which are smooth when young and slightly reticulate after maturity and which measure $45-50 \times 20-23 \mu$. The descriptions and published figures of the fungus suggest *Stemphylium botryosum* but the reticulate spore wall does not fit this species.

OTHER FORMS INCLUDED IN *STEMPHYLIUM* BUT FOREIGN
TO THIS GENUS

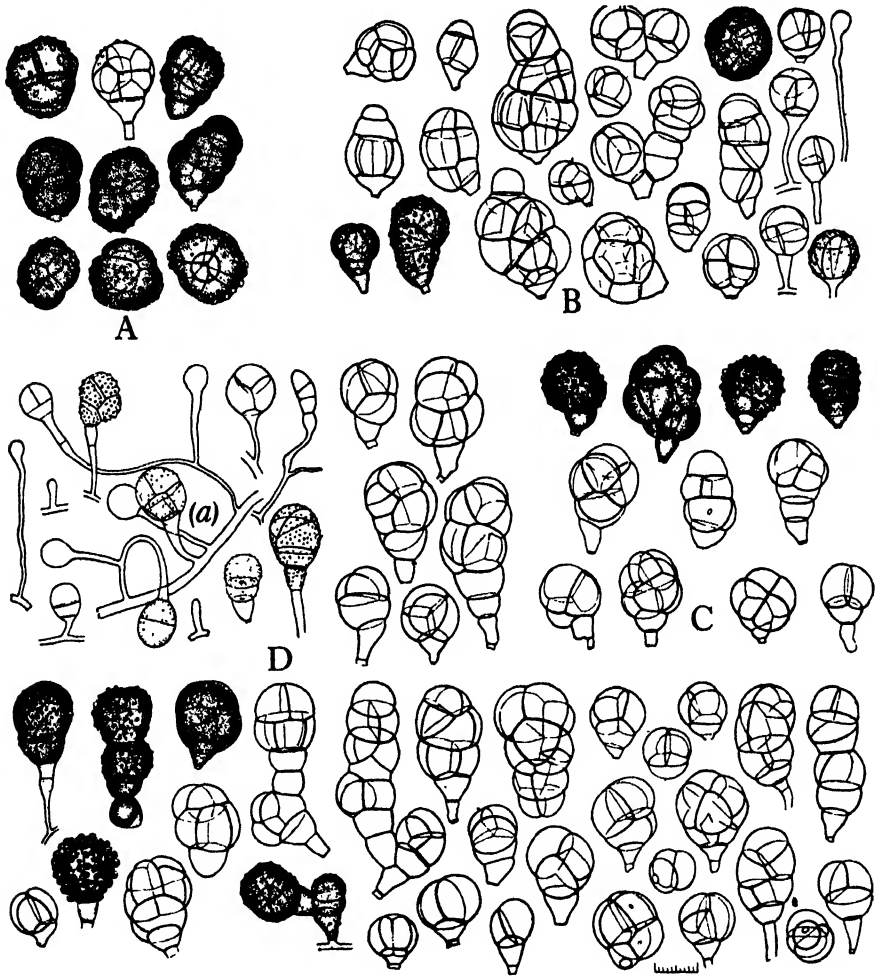
S. macrosporoideum (Berk.) Sacc.

It would be unsatisfactory to close this account of *Stemphylium* without referring to another spore form entirely distinct from the preceding ones which was introduced into this genus by Saccardo in 1886 when he transferred *Epochnium macrosporoideum* Berk. (Pl. XV, fig. 15) to the genus as *Stemphylium macrosporoideum* (Syll. iv, p. 519). This species was discovered on "the decorticated portions of a decayed twig apparently of *Ribes rubrum*" on which it formed a slate-black stratum. Berkeley's description is as follows: "From the tips or on very short lateral branches spring sub-globose or oval colourless transparent vesicles with a central nucleus; these by degrees are furnished with obscure septa still retaining their transparency; at length they acquire when full grown a brown hue and are from $\frac{1}{1500}$ to $\frac{1}{2000}$ of an inch in diameter. They are then in general more or less globose, divided by septa into a few lobes, which are disposed in a radiating manner like the berries of a mulberry. Occasionally the septa appear darker than the rest of the sporidia. A few are furnished with a little apicular peduncle but the greater part lose all traces of the point of attachment. I have sometimes seen one or two cells projecting from the otherwise globose sporidia and in one instance two sporidia were united by means of such a process." The twig comprising Berkeley's original specimen is preserved in Herb. Kew. but from it I could obtain only a very few spores. These agreed with the description given by Berkeley but the surface was distinctly warted on some of them.

In 1932 I found Berkeley's *Epochnium macrosporoideum* growing on the underside of damp linoleum in my house at Kew. The fungus produced an extensive growth, which, when uncontaminated with other fungi, was a beautiful lavender colour. The species readily grew in culture and the mode of development was exactly as described by Berkeley and the different stages are shown in Text-fig. 13. The so-called "apicular peduncle" is clearly formed by the somewhat conically shaped basal cell of the spore, which is seen projecting from most spores below which the stalk breaks off when the spore is shed. Sometimes there is a sort of lop-sided swelling to the "subglobose or oval vesicle" of the young spore but nothing in the form of a spiral is seen such as occurs in a related species. The spores measure $10-49 \times 9-18 \mu$, mostly about $19 \times 13 \mu$.

Besides Berkeley's original material, there is in Herb. Kew. a specimen labelled in Cooke's handwriting "*Stemphylium macrosporoideum* (B. & Br.) Sacc." without any indication of its origin. It

consists of the characteristically coloured growth on a piece of bark and bears a large number of spores which microscopically agree with those figured by Berkeley, especially as regards the branched spores or

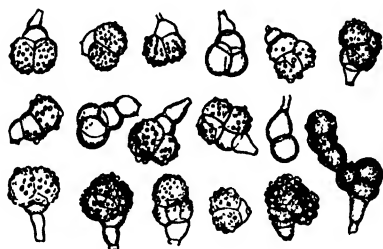


Text-fig. 13. *Acrospeira macrosporoidea* (= *Stemphylium macrosporoideum* (B. & Br.) Sacc.). A. Conidia from type material of *Epochnium macrosporoideum* B. & Br. in Herb. Kew. B. Conidia from a specimen labelled "*Stemphylium macrosporoideum* (B. & Br.) Sacc." in Cooke's handwriting in Herb. Kew. C. Conidia from a collection on linoleum in Kew, 1932. D. Conidia from an isolation from C in culture on clear maize meal and onion agars. Stages in the development of the spores are shown at (a). Magnification $\times 500$.

spores united by means of a process, examples of which were not found in the type. Similar spores were obtained in culture, where they developed just above the surface of the agar. The systematic position of this species and of the two following is discussed below.

***Stemphylium aspersorum* Cooke & Masee**

A second species of the same type as *Stemphylium macrosporoideum* is *S. aspersorum* Cooke & Masee, 1887. It was discovered on paper in London. The original specimen in Herb. Kew. agrees with the diagnosis except that the conidia themselves are about 12μ in diameter and not "each cell" of the conidium. The development of the fungus has not been followed in culture but it evidently agrees with *S. macrosporoideum* in the way the spores are borne, the stalk cell being clearly recognizable while a few immature spores seen in the type material show that the same kind of attachment obtains [Text-fig. 14].

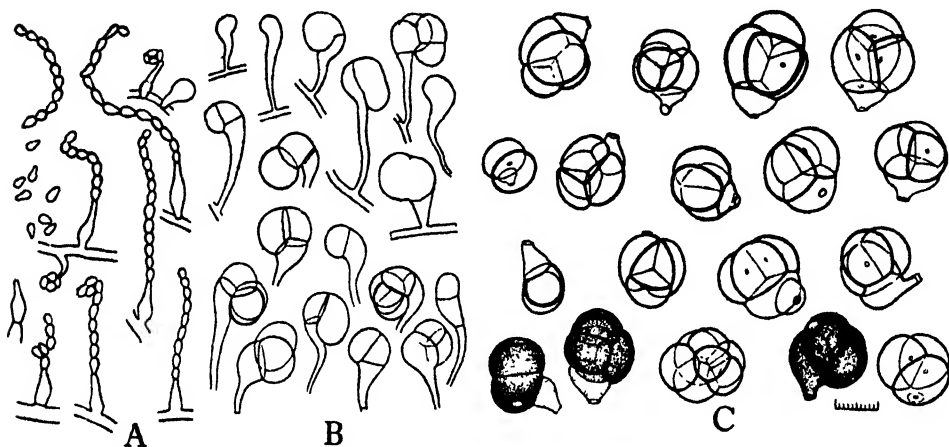


Text-fig. 14. *Acrospeira aspersora* (= *Stemphylium aspersorum* Cooke & Mass.) drawn from type material in Herb. Kew. $\times 500$.

***Stemphylium paxianum* (von Szabó) Lindau**

A third species which is allied to *Stemphylium macrosporoideum* is one received from Baarn under the name of *S. paxianum* (von Szabó) Lindau (Text-fig. 15). This species grows well in culture, forming a silver-grey colony bearing large numbers of spores. These are initiated by a swelling of the end of the hypha (or of a lateral branch), which becomes somewhat clavate but is usually more or less twisted or bent. The swelling at the end enlarges and may partly turn over to give a slight indication of a hook. The young spore is then cut off by a wall, further swelling takes place, the hook may become more marked, other walls are formed and ultimately a roughly spherical, thick-walled, dark-coloured spore is produced, divided in all planes into one to six (mostly four) cells and provided with a broadly obconical basal cell, the free end of which marks the point of attachment to the conidiophore; the broken end of the latter may remain attached to the spore-body. The surface is smooth and germ-pores can be recognized, usually one to each cell. Pores are also frequently visible in the dividing walls. The spores measure $12-25 \times 10-19\mu$. After careful search phialospores were found in old cultures which had been revived. These are obpyriform, hyaline, about $2-4 \times 1.5-2\mu$ and occur in chains of up to fourteen phialospores, being abstricted in succession from a subulate basal cell borne as a branch on the mycelium.

S. paxianum was originally described as *Tetracoccusporium paxianum* in 1905. I am indebted to Dr von Szabó for the information that no type material of the species exists, but from the diagnosis and accompanying figures it is clear that the spore was divided into four cells by two vertical walls at right angles to each other. A specimen of the Baarn fungus was submitted to Dr von Szabó, who stated that it was not his species. The Baarn culture was derived in the first instance from deer dung in Belgium. So far as I am aware the species has not yet been given a name, though it appears probable that it has been taken to be *S. macrosporoideum* by various mycologists. Oudemans's isolation of the latter species preserved at the Centraalbureau, Baarn, proved to be identical with it. The descriptions of Saccardo



Text-fig. 15. *Acrospeira levis* from the culture sent to Baarn as *Stemphylium paxianum* (von Szabó) Lindau. A. Phialospores developed in a month-old culture on clear maize meal agar. B. Stages in the development of the aleuriospores. C. Mature aleuriospores. Note the pores in the cross septa and the germ pores in two of the shaded spores. Magnification $\times 500$.

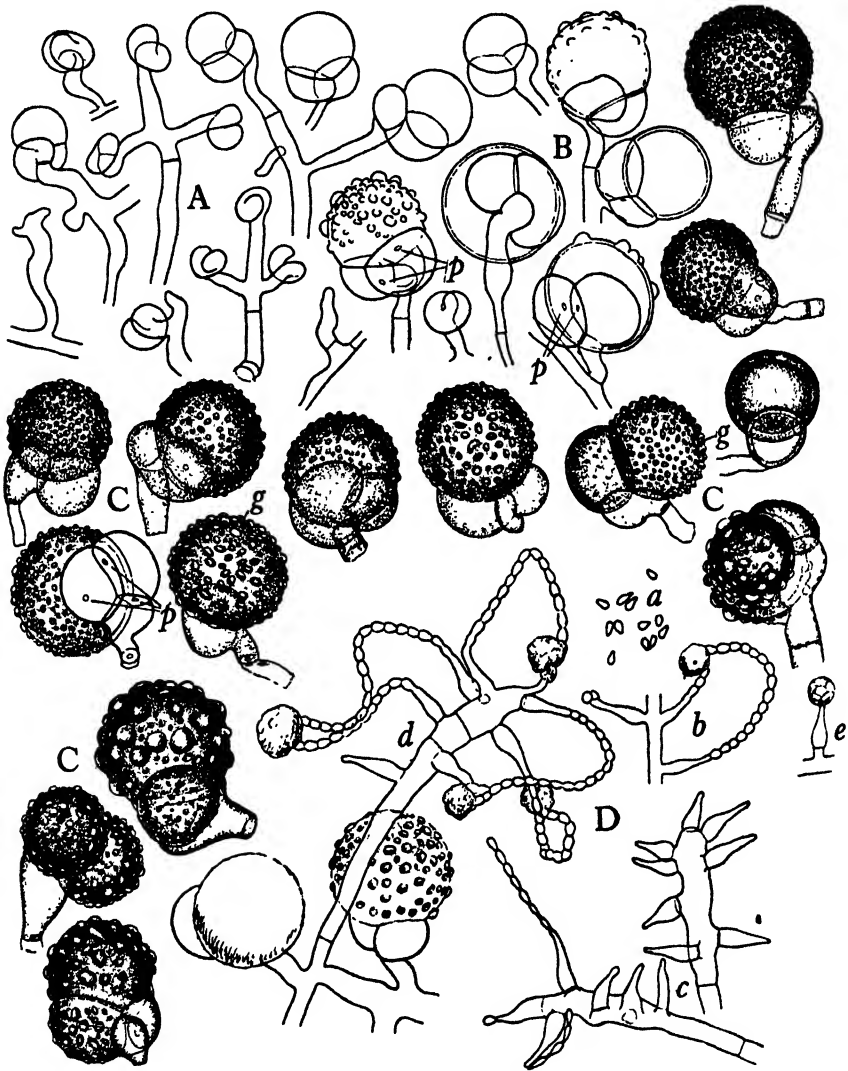
and, later, those of Lindau and Ferraris mentioned the spores of *S. macrosporoideum* as "cruciately or radially divided". Berkeley stated that the spores of his species are disposed in a radiating manner but the cruciate character is not mentioned; it may have been taken from Berkeley's figure *e*. The spores of the Baarn fungus, however, are not cruciate though those of the true *Tetracoccusporium paxianum* are and it may have been this character, erroneously attributed to *Stemphylium macrosporoideum*, which induced Lindau (1908, p. 219) to transfer *Tetracoccusporium paxianum* to *Stemphylium*.

It will probably be conceded that *S. macrosporoideum*, *S. aspersorum*, and the Baarn culture of *S. paxianum* belong to the same genus but not to *Stemphylium*. On the other hand, they have much in common with *Acrospeira*, and this genus may now be considered in this connexion.

THE GENUS *ACROSPEIRA* IN RELATION TO *STEMPHYLIUM*
MACROSPOROIDEUM AND ALLIED FORMS

Berkeley & Broome in 1861 erected *Acrospeira* for the remarkable fungus *A. mirabilis* found on Spanish chestnuts at Bristol. The structure of this fungus has been studied by Hotson (1912), by Mangin & Vincens (1920) who named it *Spirospora Castaneae* (erecting a new genus) and by Peyronel (1921) who discussed in detail the synonymy. The characteristic feature of the species is the spiral coiling of the young conidial primordia (Text-fig. 16). The ends of the branches of the conidiophore, usually three at first but finally numerous, swell slightly and become coiled, the coil often occurring in more than one plane. Transverse divisions, usually three, develop in the coiled portion, cutting off three cells, and the terminal one increases in size and becomes globular. The spiral twisting is so tight that the two lower cells lie closely adpressed to the large terminal cell which is warted and coloured brown. Hotson (p. 266), following Berkeley & Broome and Biffen, considers that it is the penultimate cell of the coil which enlarges as a rule to form the functional spore, but occasionally the end cell. In cultures very kindly supplied to me by Prof. Peyronel only the terminal cell was observed to do this and confirmatory evidence of this was obtained from the fully developed spore. Buller (1933, p. 86) has recently discussed the pores which occur in cross-walls of fungus hyphae. In the mature spore of *Acrospeira mirabilis* such pores can be recognized in some of the walls dividing the cells. If the enlarged cell were the central cell of the three cells constituting the young spore, then the walls between it and each of the two smaller cells should have pores, and there should be no pore in the wall between the two small cells. On the other hand, if the enlarged cell is the terminal cell of the coil, there should be a pore only in one of the walls dividing the large cell from the two smaller cells and a pore in the wall dividing the two small cells from each other. The latter alternative has been repeatedly observed and is represented in Text-fig. 16*p*, but no instance of the former has been seen. The number of cells constituting the spore may vary from two to four or more but only the terminal cell swells up; the stalk cell is usually clearly evident, sharply marked off by its brown colour from the supporting hyaline hypha. The spore is shed apparently by being broken off. A single germ pore (Text-fig. 16*g*) is often recognizable on the enlarged cell at a point diametrically opposed to the perforated cell wall dividing it from the next cell below and the single germ-tube usually develops radially from the pore (see Mangin & Vincens, Fig. 7). The straight germ-tube branches near its base, pushing the spore aside, so that the branch comes to lie in the same straight line as the original germ-tube, but growing in an opposite direction. Mangin & Vincens described

the phialospore stage of the fungus and this was readily obtained in pure culture on potato dextrose agar. The phialospores are obpyriform, borne in long chains, arising from phialids formed directly on



Text-fig. 16. *Acrospeira mirabilis* B. & Br. isolated from chestnut by Prof. Peyronel. Drawing from cultures on various media showing: A. Young stages in the formation of the aleuriospores. B. Intermediate stages. C. Fully developed aleuriospores. D. Phialospores. a, spores in lactic acid; b-e, phialospore development (from living cultures in surface view; d shows phialospores and aleuriospores on the same hypha). Note septal pores at p and germ pores at g. $\times 500$

the mycelium or on the swollen end of a hypha projecting into the air.

We are now in a position to consider the relation between *Acrospeira* and *Stemphylium macrosporoideum* and allied species. In all these species

the spores consist of a number of cells united to form a muriform, globose spore, the stalk cell of which is recognizable as a more or less obconical projection. The spore does not fall from the conidiophore leaving a scar but is broken off and often the remnants of the fractured hypha remain attached. This behaviour marks off these species from *Stemphylium* as here interpreted. The method of spore development in these species, however, is not uniform; in *Acrospeira* the spore develops as a closely pressed coil; in the Baarn "*Stemphylium paxianum*" there is some suggestion of a hook but cells of the spore are formed by subdivision of the "hook" rather than by the coiling of the transversely septate spore primordium; in *S. macrosporoideum* the young spore is almost spherical and the cells arise by subdivision without any coiling, while in *S. asperosporum* it is possibly similar. Further, the phialospores of *Acrospeira mirabilis* and the Baarn "*Stemphylium paxianum*" are of the same general type; they have not been observed in the other two species. *Acrospeira* has usually been regarded as belonging to the Dematiaceae-Amerosporae (see Shear and Clements, Ferraris) but the spore clearly consists of more than one cell, and as the walls dividing the cells are oblique, it may more properly be placed in the Phaeodictyae. Hotson (p. 298) remarked on the similarity between the mature spores of *Stemphylium macrosporoideum* and *Acrospeira mirabilis* (though he regarded the spores of the latter as bulbils). Apart from the method of spore development, and also the colour of the spores, which is reddish brown in *A. mirabilis* and olivaceous in the others, there is considerable similarity between all these species and the question arises whether they are sufficiently alike to be regarded as belonging to one genus. There are two alternatives: (1) to erect a new genus for *Stemphylium macrosporoideum*, *S. asperosporum* and the Baarn "*S. paxianum*"; (2) to transfer all these species to *Acrospeira*.¹ The multiplication of genera in the present stage of systematic mycology, will, if allowed to proceed unchecked, ultimately result in a large number of monotypic genera being established, the relationships of which will remain obscure. A better plan is to group more or less similar species round known generic types, leaving the limits of the genera to become manifest in course of time. Furthermore, many of the genera are not properly understood and to add to their number

¹ Montagne in *Ann. Sci. nat. Bot. Sér. iv*, viii, 299 (1857), erected a genus *Acrospeira* on the single species *A. Crouanii* found on *Angelica sylvestris*. Saccardo remarks (*Syll.* xiv, p. 1056) that Montagne's genus is related to, but distinct from, *Acrospeira* B. & Br., whilst Clements and Shear regard it as a doubtful genus. Berkeley briefly referred to *A. mirabilis* in his *Introduction to Cryptogamic Botany* (p. 305 and Fig. 96a), 1857, but this description probably cannot pass as a valid generic diagnosis. Furthermore there is some doubt whether the *Introduction* was published earlier in 1857 than Montagne's genus, or not. It is possible that the name *Acrospeira* B. & Br. may prove to be untenable, but as it is so well established I think it best to adhere to it for the present, especially as the spelling is different from Montagne's.

without a very real necessity is to be deprecated. The chief objection to the second alternative is that the characteristic method of spore development in *A. mirabilis* is not followed by the other species but the Baarn "*Stemphylium paxianum*" forms a connecting link which greatly strengthens the case for uniting all the species in the one genus. Therefore, while admittedly widening the conception of *Acrospeira* rather more than some may care to do, I propose to adopt what seems to me to be the lesser of two evils and to transfer the species *Stemphylium macrosporoideum* and *S. asperosporum* to *Acrospeira* as *A. macrosporoidea* (Berke) comb.nov. and *A. asperospora* (Cooke & Massee) comb.nov. respectively and to name the Baarn culture of "*Stemphylium paxianum*" *Acrospeira levis*. This, at any rate, will be more satisfactory than leaving these species in *Stemphylium*, where they are manifestly out of place.

The diagnosis of *Acrospeira levis*¹ is as follows:

Acrospeira levis sp.nov.

Hyphis septatis, hyalinis, $3\ \mu$ crassis, numerosissimas aleuriosporas gerentibus, et colonias argenteas cinereas formantibus.

Aleuriosporis solitariis, hyphis principibus, vel ramis brevibus insidentibus, initio clavatis, dein apice turgescitibus, sinuatis, et aliquando recurvatis, in 2-6 (saepè 4) loculos clathratos partitis, fere globosis, loculo inferiori obconico praeditis, $12-25 \times 10-19\ \mu$; episporio levi, crasso, fuligineo ad septa constricto.

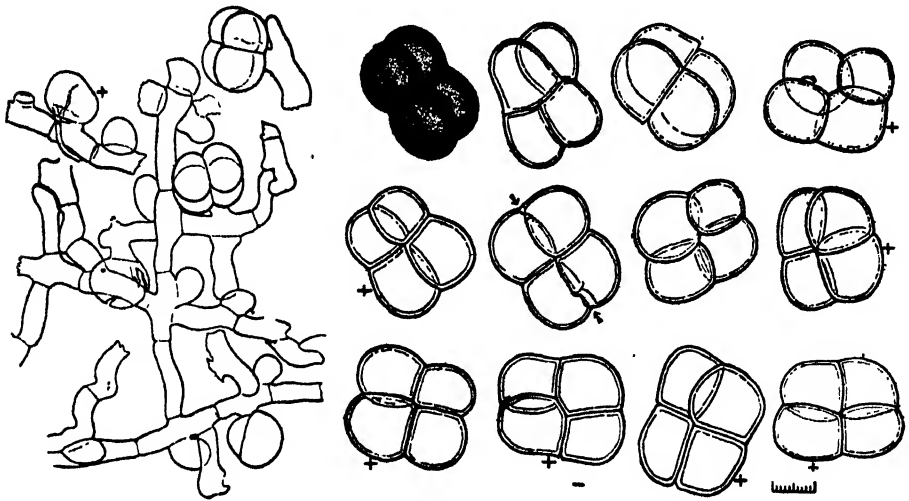
Phialidis raris, hyalinis, solitariis, subulatis, prope basim 1 septatis, $6-11 \times 2-3.5\ \mu$; phialosporis in catenulas (2-14 sporas) vel mucosa capitula abstrictis, continuis, hyalinis, obpyriformibus, $2-4 \times 1.5-2\ \mu$.

Stemphylium quadratum (Cooke) Sacc.

Finally the species *S. quadratum* needs to be considered. Originally described as *Epochnium quadratum* by Cooke in 1883, it was renamed *Stemphylium quadratum* by Saccardo in 1886 (*Syll.* iv, 521), and original material of Cooke's preserved in Herb. Kew. shows thick-walled, fragile, very dark-coloured spores of four cells arranged mostly cruciately (Text-fig. 17). From the material it is difficult to make out how the spores are borne but they seem to be attached near to the junction of the vertical walls; and to arise as lateral swellings on the mycelium. Such a spore structure cannot be included in *Stemphylium* as here revised. The genus *Tetracoccusporium* from the original description and figure (as mentioned on p. 232 no type material exists) was evidently based on a fungus with similar spores and Prof. Dr von Szabó, who kindly examined a drawing of the spores of *Epochnium quadratum* Cooke (Text-fig. 17) was of opinion that the latter species

¹ I am indebted to Mr E. W. Mason for help with the Latin.

was closely related to *Tetracoccusporium paxianum*. The spore measurements, however, are larger and I accordingly adopt the genus *Tetracoccusporium* and transfer *Stemphylium quadratum* to it as ***Tetracoccusporium quadratum*** (Cooke) comb.nov., a repetitive name it is possible to avoid.



Text-fig. 17. *Tetracoccusporium quadratum* (= *Stemphylium quadratum* (Cooke) Sacc.) drawn from type material of *Epochnium quadratum* Cooke on *Fraxinus* (labelled No. 3338, not 3388 as in *Grevillea*) in Herb. Kew. except spores indicated by +, which are from the authentic exsiccatum Rav. Amer. Fungi 773. A. Showing the formation of spores on the superficial mycelium. B. Mature spores. Fractures of the wall of one spore are indicated by arrows. Magnification $\times 500$.

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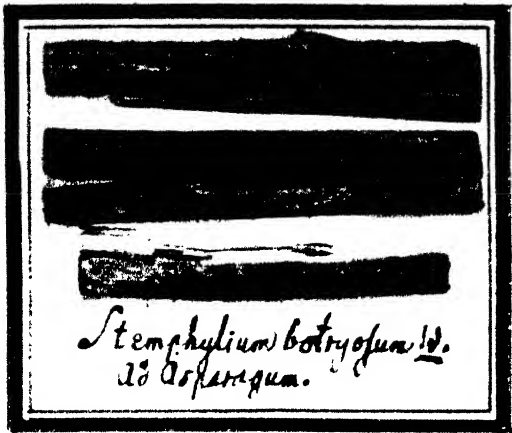


Fig. 1



Macrosporum varicella

Fig. 2



Fig. 3



Fig. 4



Fig. 5

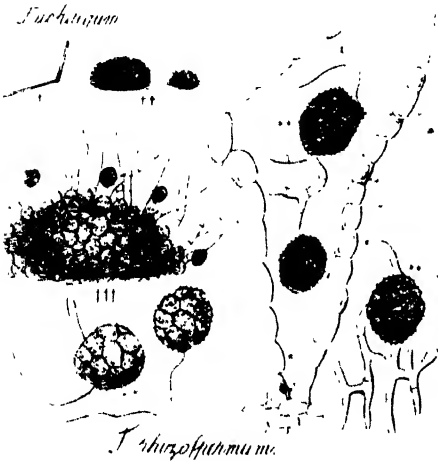


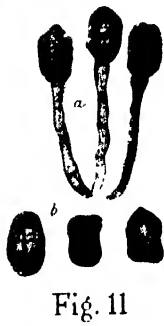
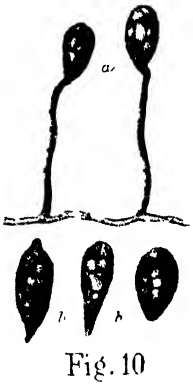
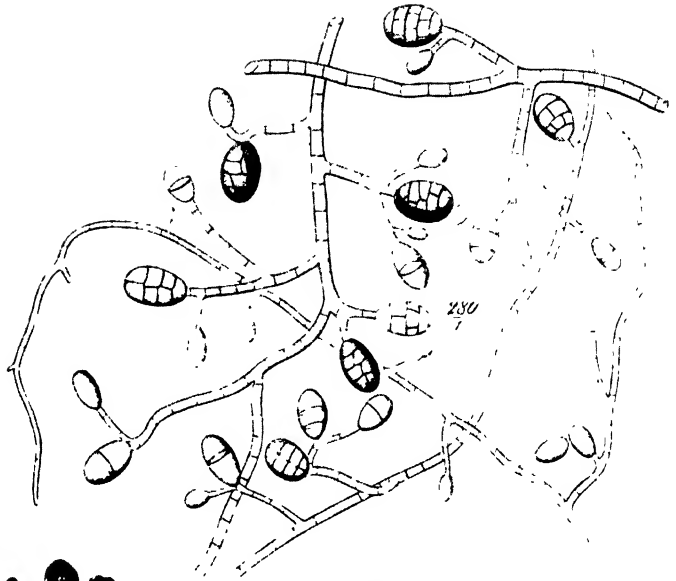
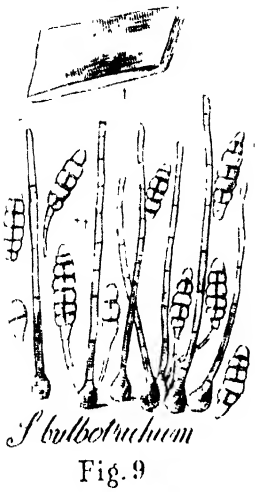
Fig. 6



Fig. 7



Fig. 8



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EXPLANATION OF PLATES XIV AND XV

PLATE XIV

- Fig. 1. *Stemphylium botryosum* Wallr. Type specimen. Natural size.
- Fig. 2. *Macrosporium sarcinula* Berk. Berkeley's original figure.
- Fig. 3. *Stemphylium pyriforme* (Cda) Bon. (= *Sporidesmium pyriforme* Cda).
- Fig. 4. *Stemphylium polymorphum* (Cda) Bon. (= *Sporidesmium polymorphum* Cda).
- Fig. 5. *Stemphylium elegans* (Cda) Bon. (= *Sporidesmium elegans* Cda).
- Fig. 6. *Stemphylium rhizospermum* (Cda) Bon. (= *Trichaegum rhizospermum* Cda).
- Fig. 7. *Stemphylium graminis* (Cda) Bon. (= *Soredospora graminis* Cda).
- Fig. 8. *Stemphylium dubium* (Cda) Bon. (= *Mystrosporium dubium* Cda).

Figs. 3-8. Corda's original figures.

PLATE XV

- Fig. 9. *Stemphylium bulbotrichum* (Cda) Bon. (= *Septosporium bulbotrichum* Cda). Corda's original figure.
- Fig. 10. *Stemphylium pyriforme*. Bonorden's figure.
- Fig. 11. *Stemphylium polymorphum*. Bonorden's figure.
- Fig. 12. *Stemphylium paradoxum* (Cda) Fuck. (= *Sporidesmium paradoxum* Cda). Corda's original figure.
- Fig. 13. *Stemphylium lanuginosum* Harz. Harz's figure.
- Fig. 14. *Stemphylium sarcinaeforme* (= *Macrosporium sarcinaeforme* Cav.) on clover leaves collected at Oakley, Hants., in September, 1932. Natural size.
- Fig. 15. *Acrospeira macrosporoidea* (= *Epochnium macrosporoideum* Berk.). Berkeley's original figure.

UROCYSTIS SOROSPORIOIDES, A NEW RECORD FOR INDIA

By B. B. MUNDKUR

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(With Plate XVI)

LEAVES of a perennial species of *Delphinium* showing swellings on the blade and petiole were collected at Simla in July 1935. The species was either *D. denudatum* Wall. or *D. vestitum* Wall., but this could not be ascertained with certainty for the plants did not bear flowers at the time. Microscopic examination showed that the swellings were the pustules caused by a species of *Urocystis*.

In cross-sections the diseased parts of the blade or the petiole show an erumpent circular form (Plate XVI, figs. 1, 2, 3), the upper and lower tissues being arched outwards. The cells of the ventral side of the leaf-blade are elongated and the mesophyll is replaced by a nutrient mycelium and this, later, is apparently converted into spore masses.

When seen in mass the spores have a black appearance but the sori whilst still covered by the epidermis are greyish. The spore-balls are round, oblong or obtuse, compact and opaque, $19-51 \times 24-61 \mu$ in diameter with an average of $29 \times 38 \mu$ and consist of three to seven main fertile cells surrounded by a layer of light-coloured secondary sterile cells. The fertile cells are spherical or hemispherical, $11-16 \mu$ in diameter, dark brown with smooth episporous. The secondary cells are hemispherical, $8-12 \mu$ in diameter with yellowish brown, sub-hyaline episporous. They project slightly beyond the periphery of the spore-balls and are uniformly distributed so that the spore-balls have a fairly smooth contour. Spore-balls that are not fully mature are not always completely covered by the layer of sterile cells.

Three leaf smuts belonging to the genus *Urocystis* are known to attack members of the family Ranunculaceae. None of these has, however, been recorded for India. Of these *U. Anemones* (Pers.) Wint. has, according to Clinton (1904), spore-balls that are not completely surrounded by the secondary cells. The number of fertile cells is never more than four, while the size of the pustules is also much smaller than that of the other two smuts. *U. carcinodes* (B. & C.) de Waldh. rarely attacks the leaf-blades, according to the same authority, and is confined to the midrib, petiole, and stem. *U. sorosporioides* Koernicke,

the third smut attacking this family, has large sori and correspondingly large spore-balls. These latter have several fertile cells as compared with *U. Anemones* and the covering of the sterile cells is complete when mature. It attacks both the leaf-blades and petioles and causes no unusual distortion of the host. In Europe and America *U. sorosporioides* has been reported on the genus *Delphinium*, while the other two smuts have not been known to occur on this genus.

The Simla fungus agrees closely with this smut. When the spore-balls from Sydows exsicc. (No. 375) of *U. sorosporioides* were compared with those from the Simla smut, they were hardly distinguishable from them. The comparative measurements are given in Table I.

Table I. *Comparative spore-ball and spore measurements of Urocystis sorosporioides*

	Winter (1884) μ	Clinton (1904) μ	Lindau (1912) μ	Mundkur μ
Spore-balls	22-44	30-60	22-48 × 15-31	24-61 × 19-51
Fertile cells	12-17	13-17	11-17	11-16
Sterile cells	—	8-15	7-12	8-12

The measurements are in agreement to a considerable extent. The number of fertile cells in the Indian specimen is three to seven and this is in accord with the number given by the other investigators.

In the genus *Urocystis* and the older genus *Tuburcinia* the spore—more commonly called the spore-ball—is not a single cell but a globose ball consisting of a number of cells permanently bound together. In *Tuburcinia* all of these are capable of germination while in *Urocystis* only the central cells are fertile, the peripheral ones being sterile. Liro (1922), however, maintains that no such distinction exists and that a covering layer of sterile cells is common to all these species. He therefore suggests that the two genera should be merged. But Brefeld (1895), who succeeded in germinating the spore-balls of a *Tuburcinia* (*T. primulicola* Brefeld), found that the peripheral cells did germinate in large numbers and that there were no sterile cells outside of those germinated. In the spore-balls of *Urocystis*, however, there was always a persistent layer of sterile cells. As the spore-balls of only a single species of *Tuburcinia* and of a few species of *Urocystis* have so far been germinated, until the germination of the spore-balls of more of these has been studied and until there is some concord among mycologists as to the constancy of the sterile layer of covering cells, it is better to retain the genus *Urocystis*.

The smut collected at Simla on the leaves of *Delphinium* is therefore *Urocystis sorosporioides* Koernicke. Leaves of a *Delphinium* sp. collected at Chakratha near Dehra Dun and brought to Pusa for identification by Dr K. Bagchee were also found to be attacked by this same smut.

SUMMARY

A leaf smut on the leaves and petioles of a species of *Delphinium* collected at Simla in July 1935 has been identified as *Urocystis sorosporioides* Koernicke. This is a first record of this fungus for India.

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EXPLANATION OF PLATE XVI

- Fig. 1. Leaf-blade of *Delphinium* sp. showing large sori caused by *U. sorosporioides*.
Fig. 2. Petiole showing erumpent pustules of the same smut.
Fig. 3. Cross section of the petiole showing sorus and the spore-balls of *U. sorosporioides*.
× 50.



Fig. 1



Fig. 2



Fig. 3

BRITISH HYPOCREALES

By T. PETCH

(With 39 Text-figures)

THE Hypocreales constitute one of the families of the order Pyrenomycetae, ascigerous fungi in which the asci are contained in perithecia, globose, pear-shaped, or flask-shaped receptacles, furnished with a definite wall. They are distinguished from other Pyrenomycetae by the colour and consistency of their perithecia, which are brightly coloured, usually red or yellow, not black, and soft and fleshy, or membranous, never carbonaceous. The perithecia may be free and separate from one another (Nectriaceae), or they may be embedded in a brightly coloured fleshy mass, known as a stroma (Hypocreaceae). Many of them have a conidial stage which belongs to one or other of the genera of Hyphomycetae, e.g. *Fusarium*, *Isaria*, *Tubercularia*, *Dendrodochium*, etc. Much work remains to be done in the determination of the conidial stages of the various species.

The family includes several plant pathogens of prime importance. Among these are *Dialonectria galligena*, the cause of "canker" in apple trees, and *Gibberella Zeae*, the cause of "scab" of wheat and maize. It also includes a number of fungi, formerly classed together as *Hyphomyces*, which are parasitic on Basidiomycetae, and the fungi, known as *Cordyceps*, which are parasitic on insects.

No complete account of the British Hypocreales has been published since the issue in 1871 of Cooke's *Handbook of British Fungi*, in which he enumerated 56 species. Since then, many other species have been added to the British list in scattered records, often by name only, so that, unless a mycologist has access to a well-equipped mycological library, he is unable to ascertain what they indicate. Moreover, on the continent of Europe and in America, intensive researches on this group by eminent mycologists, particularly by von Höhnelt and Weese, have extended considerably our knowledge of the family, and have entailed numerous corrections and alterations both in nomenclature and classification, most of which work is inaccessible to British mycologists in general. In these circumstances, the need of a new list of British Hypocreales became urgent, and the present revision was undertaken, by request, with the object of bringing an up-to-date list within the reach of working mycologists, and providing them with a means of identification of the species. It may be noted that, owing to the lack of such a list, several important plant pathogens have become generally known by names which really belong to saprophytic species.

Modern genera have been adopted wherever possible, in the belief that identification is facilitated by division into smaller groups. In the genus *Nectria*, the largest in the family, Saccardo's subdivision has been followed, though it has not been permissible to use his names in all cases. Cooke, in *Grevillea*, XII, raised some of Saccardo's subgeneric names to generic rank, and established the division of the genus *Nectria* into species on a fleshy stroma, *Nectria*, and non-stromatic species, *Dialonectria* (Greek, *dialuo*, to separate). Seaver, in *Mycologia*, I (1909), 42, though maintaining Cooke's division, transferred the name *Nectria* to Cooke's *Dialonectria* and proposed a new name, *Creonectria*, for Cooke's *Nectria*. Seaver's alteration, however, was based on a misapprehension and cannot be accepted.

Further extensive changes have been made in the genus *Hypomyces*, which, as instituted by Tulasne, was a biological genus, including those species which are parasitic on Basidiomycetae. Its constituent species have been split off at different times by different authors, and the British species now stand in four genera.

Reasons for rejecting several names and reducing others to synonymy have been published in the *Journal of Botany*, LXXIII (1935), 184-92, 217-24; LXXIV (1936), 185-93; LXXV (1937), 217-31. A few species remain doubtful, as no specimens are known; in such cases, the original description has been quoted. Although many of the older names have had to be discarded, the list now contains 124 species distributed in 41 genera.

I tender my thanks to Mr J. Ramsbottom, Keeper of Botany, British Museum (Natural History), and to Miss E. M. Wakefield, Royal Botanic Gardens, Kew, for their assistance in the examination of the Hypocreales in those herbaria, and to Dr C. G. C. Chesters, Botanical Department, Birmingham University, for similar assistance as regards the Plowright Herbarium.

HYPOCREALES

Perithecia bright-coloured, red, yellow, violet, or pallid, opening by a circular pore (ostium), immersed in the host, or superficial, separate, scattered, or caespitose on a superficial stroma, or embedded in a superficial or immersed stroma; perithecial wall membranous or fleshy, never carbonaceous; stroma, when present, bright coloured, fleshy, more or less soft, or byssoid.

Family 1. NECTRIACEAE

Perithecia separate, immersed in the host or superficial, or crowded on a superficial fleshy and parenchymatous stroma, or on a byssoid subiculum.

Family 2. HYPOCREACEAE

Perithecia immersed in a stroma, which is bright coloured, fleshy and parenchymatous, superficial or immersed in the host.

NECTRIACEAE

A. Ascospores continuous, hyaline, not filiform.

- | | |
|--|----------------------|
| Perithecia immersed in the host, ascospores oval | <i>Hyponectria</i> |
| Perithecia immersed in the host, ascospores spherical | <i>Battarrina</i> |
| Perithecia superficial, on a continuous byssoid subiculum | <i>Byssonectria</i> |
| Perithecia superficial, without a continuous subiculum or stroma | <i>Pseudonectria</i> |

B. Ascospores continuous, fuscous, dark brown or black.

- | | |
|--|---------------------|
| Perithecia superficial, with a long beak, or with setae round the ostium | <i>Melanospora</i> |
| Perithecia superficial, without a beak, or setae round the ostium | <i>Sphaeroderma</i> |

C. Ascospores one-septate, hyaline, not ciliate.

- | | |
|--|----------------------|
| Perithecia caespitose | |
| on an erumpent or superficial stroma | <i>Nectria</i> |
| at the base of a <i>Stilbella</i> conidial stage | <i>Sphaerostilbe</i> |
| Perithecia superficial, scattered or crowded | |
| without a continuous subiculum | |
| perithecia smooth | <i>Dialonectria</i> |
| perithecia hairy | <i>Lasionectria</i> |
| perithecia bearing long triangular processes | <i>Neohenningsia</i> |
| on a continuous byssoid subiculum | |
| ascospores more or less obtuse | <i>Hyphonectria</i> |
| ascospores apiculate | <i>Hypomyces</i> |
| ascospores unequally septate | <i>Apiocrea</i> |
| ascospores dividing into two | <i>Protocrea</i> |
| Perithecia immersed in the host, then erumpent | <i>Nectriella</i> |

D. Ascospores one-septate, hyaline, ciliate.

- | | |
|--------------------------------------|-----------------------|
| Ascospores with a cilium at each end | <i>Rhynchonectria</i> |
|--------------------------------------|-----------------------|

E. Ascospores one-septate, brown.

- | | |
|------------------------|-------------------|
| Perithecia superficial | <i>Letendraea</i> |
|------------------------|-------------------|

F. Ascospores oblong, fusoid, or cylindric, two- or more septate.

- | | |
|--|--------------------|
| Perithecia superficial, without apical teeth, ascospores not ciliate | |
| wall red, yellow or pallid | <i>Calonectria</i> |
| wall violet or blue by transmitted light | <i>Gibberella</i> |
| Perithecia superficial, with spreading teeth at the apex | <i>Actiniopsis</i> |

G. Ascospores filiform, or elongated fusoid, hyaline.

Torrubiella

Trailia

Pleonectria

Selinia

coloured, stroma horizontal *Chromocrea*

ascospores continuous *Claviceps*

HYPONECTRIA Sacc. in *Michelia*, I (1878), 250

Hyponectria Buxi (Desm.) Sacc. in *Michelia*, 1 (1878), 250; *Sphaeria Buxi* Desm. in *Ann. Sci. Nat.* ser. 2, XIX (1843), 354; *Sphaerella Buxi* (Desm.) Fuckel, *Symb. Mycol.* (1869), 100; *Sphaerella Buxi* (DC.) Cooke (in part), *Handbook* (1871), 922; *Sphaerella Buxi* (Desm.)

Auersw., *Syn. Pyrenom.* (1869), 2; *Sphaerella* (*Laestadia*) *Buxi* Fuckel, Cooke in *Journ. Bot.* (1883), 68; *Laestadia Buxi* (Fuckel) Sacc., *Syll. Fung.* II (1883), xxxi; *Trochila Buxi* Capron, in Cooke, *Handbook* (1871), 768.

Perithecia hypophyllous, immersed, each in a small, rose-coloured, depressed spot, which may become brown when old, 0.36 mm. diameter, depressed globose; wall of perithecium yellow; epidermis turning brown over the apex of the perithecium; asci oblongo-clavate, $66 \times 11 \mu$, spores biseriate; ascospores narrow-oval or fusoid, ends rounded, sometimes inequilateral, sometimes with the lower end subtruncate, hyaline, continuous, $14-18 \times 4-5 \mu$.

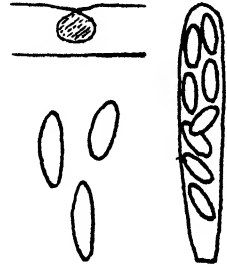


Fig. 1. *Hyponectria Buxi*; perithecium in leaf, $\times 40$; ascus, $\times 400$; ascospores, $\times 500$.

On leaves of Box. Milton, Northants (Berkeley); Forden (Vize), in Plowright, *Sphaer. Brit.* no. 8, and Cooke, *Fung. Brit. Exsicc.* II, no. 478; Sandsend, Yorks, Hb. B.M.; Overstrand Woods, Norfolk, October 1934; Helmsley, Yorks, August 1935.

BATTARRINA Sacc., *Syll. Fung.* II (1883), 533 (as subgenus)

Perithecia immersed in the hymenium of the host (*Tuber puberulum*), globose, hyaline, sometimes confluent; asci cylindrical, eight-spored; ascospores hyaline, continuous, globose.

Battarrina inclusa (B. & Br.) Sacc., *Syll. Fung.* II. (1883), 533; *Hypocrea inclusa* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 3, VII (1861), 451, pl. 17, fig. 23.

Perithecia globose, about 0.25 mm. diameter, hyaline, sometimes confluent, astomate; wall thin, of parallel hyphae up to 4μ diameter; asci at first broadly clavate, $16-20 \times 5-6 \mu$, eight-spored, spores biseriate, becoming cylindric, $30 \times 5 \mu$, soon diffluent, spores uniseriate, persisting in lines; ascospores globose, hyaline, $4-5 \mu$ diameter, ornamented with an interrupted network of narrow bands which form a narrow border to the spore.

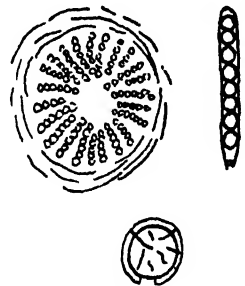


Fig. 2. *Battarrina inclusa*; cross-section of perithecium, $\times 100$; ascus, $\times 1000$; ascospore, $\times 1500$.

Leigh Wood, Bristol, September–October 1859 (Broome), Hb. Kew. and Hb. B.M.; Hanham near Bristol, November 1869, November 1871 (Broome), Hb. Kew. and Hb. B.M., Rabh. *Fung. Europ.* no. 1610.

BYSSONECTRIA Karst. in *Medd. Soc. Fauna Fl. Fenn.* VI (1881), 6

Perithecia separate, superficial, on a common byssoid subiculum, bright-coloured, soft; asci eight-spored; ascospores hyaline, continuous, not filiform.

Byssonectria viridis (A. & S.) Petch in *Journ. Bot.* LXXV (1937), 220; *Sphaeria viridis* A. & S., *Consp. Fung.* (1805), 8, pl. 6, fig. 8; *Sphaeria luteovirens* b. Fries, *Syst. Mycol.* II (1822), 339; *Hypomyces viridis* (A. & S.) B. & Br. in *Ann. Mag. Nat. Hist.* ser. 3, xv (1865), 451; *Hypomyces luteovirens* Plowr. in *Grevillea*, XI (1882), 46; *Peckiella viridis* (A. & S.) Sacc., *Syll. Fung.* IX (1891), 944; *Hypomyces ater* Cooke in *Grevillea*, XII (1884), 80; *Peckiella atra* (Fr.) Sacc., *Syll. Fung.* IX (1891), 944.

Subiculum thin, at first bright egg-yellow, becoming greenish and then almost black; perithecia crowded, ovoid, pallid, becoming brown or nearly black, 0.38 mm. high, 0.3 mm. diameter; asci cylindrical, $160 \times 7-8 \mu$; ascospores narrow-oval or fusoid, sometimes inequilateral, ends produced into rather long, solid tips, which are often curved or hook-shaped, continuous, minutely warted, $27-45 \times 5-6 \mu$.

Conidia oval, hyaline (Plowright).

On various agarics, especially *Lactarii*. Northamptonshire, July 1848, Hb. Kew. and Hb. B.M. South Wootton, 1878-9-80 (Plowright), Hb. B.M., Rehm, *Ascomyceten*, no. 586; Carlisle (Cooke), Hb. Kew.; Forres, September 1896 (Keith), Hb. Kew.

Byssonectria lateritia (Fr.) Petch in *Journ. Bot.* LXXV (1937), 220; *Sphaeria lateritia* Fr., Kunze, *Myc. Hefte*, II (1823), 42; *Hypocrea lateritia* Fr., *Summa Veg. Scand.* (1849), 383; *Hypomyces lateritius* Tul. in *Ann. Sci. Nat.* ser. 4, XIII (1860), 11; *Peckiella lateritia* (Fr.) Maire in *Ann. Mycol.* IV (1906), 331.

Subiculum white, becoming pale yellow or yellowish brown; hyphae stout, with inflated segments, $15-18 \times 7-9 \mu$, the terminal segment conoid, or broadly flask-shaped, or ovate, $15-20 \times 9-10 \mu$, moderately thick-walled, pale brown; perithecia semi-immersed, ovato-globose, 0.3 mm. high, 0.25 mm. diameter, apex papillate, yellowish or brownish, darker than the subiculum, wall hyaline, becoming brown; asci cylindrical, with a long pedicel, $125-160 \times 5-9 \mu$, spores uniseriate or obliquely uniseriate; ascospores lanceolate or narrow-oval, sometimes inequilateral, apiculate, sometimes acuminate, minutely verrucose, continuous, hyaline, $15-25 \times 4-5 \mu$.

Conidia globose, hyaline, $3.5-7 \mu$ diameter, on subulate conidiophores, $30-40 \mu$ long (Tulasne).

On *Lactarii*, especially *L. deliciosus*. Leigh Wood, September 1844 (Broome), Hb. Kew. and Hb. B.M.; Leigh Wood, October, November, 1860 (Broome), Hb. Kew. and Hb. B.M., Rabh. *Fung. Europ.* no. 317; Hereford, 1871-74, Hb. Kew. and Hb. B.M., Plowright, *Sphaer. Brit.* I, no. 5; King's Lynn, October 1882 (Plowright), Hb. Kew., de Thümen, *Mycoth. Univ.* no. 2164; etc.

The specimens attributed to *Hypomyces torinosus* (Mont.) Tul., on *Lactarius*

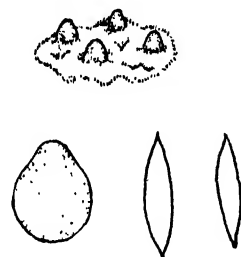


Fig. 3. *Byssonectria lateritia*; group of perithecia, $\times 15$; perithecium, $\times 50$; ascospores, $\times 750$.

terminosus, Dinmore, October 1874, Plowright, *Sphaer. Brit.* II, no. 4, Cooke, *Fung. Brit. Exsicc.* II, no. 667, Vize, *Microf.* no. 587, are *Hypomyces lateritius*, as are also "*Hypomyces terminosus*", North Wootton, October 1896 (Plowright), Hb. Kew., and King's Lynn, October 1900 (Plowright), Hb. B.M.

PSEUDONECTRIA Seaver in *Mycologia*, I (1909), 48

Perithecia separate, superficial, without a common subiculum, bright-coloured, soft; asci eight-spored; ascospores hyaline, continuous, not filiform.

Pseudonectria Rousseliana (Mont.) Seaver in *Mycologia*, I (1909), 48; *Nectria Rousseliana* Mont., *Syll. Crypt.* (1856), 224; *Nectriella Rousseliana* (Mont.) Sacc. in *Michelia*, I (1877), 51.

Perithecia superficial, gregarious, subglobose or conoid, 0.14–0.3 mm. diameter, 0.16–0.3 mm. high, straw-coloured, or brick-red, or greenish hyaline, sparsely clothed with hyaline, rigid, spreading setae; setae up to 100 μ long, 5 μ diameter, continuous or sparingly septate, apex obtuse or subacute; asci clavate, attenuated below, 54–60 \times 6–8 μ , apex truncate, spores obliquely uniseriate, then biseriate; ascospores oval or somewhat lozenge-shaped, hyaline, continuous, thick-walled, 9–16 \times 3–5 μ .

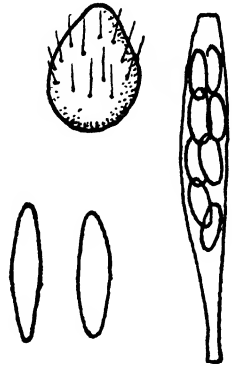


Fig. 4. *Pseudonectria Rousseliana*; perithecium, $\times 50$; ascus, $\times 650$; ascospores, $\times 1000$.

On leaves of Box. Milton, Northants (Berkeley), Hb. Kew. Twycross, November 1856 (Bloxam), Hb. B.M. and Hb. Kew.; Batheaston, February 1859 (Broome), Hb. B.M. and Hb. Kew.; Elmhurst, March 1859, Hb. B.M.; Spye Park (Broome), Hb. Kew.; Dorking, Cooke, *Fung. Brit. Exsicc.* no. 597; Scarborough (Masse), Hb. B.M.; Kilnwick Percy, Yorks, August 1936.

Pseudonectria furfurella (B. & Br.) Petch, comb. nov.; *Nectria furfurella* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 4, VII (1871), 435; *Nectriella furfurella* (B. & Br.) Sacc. in *Michelia*, I (1878), 278.

"Perithecia on an effused fleshy stroma, flesh-coloured, subglobose, collapsing, beset with glistening scale-like particles. Paraphyses branched; sporidia ovate, 3.75–5 μ long; conidia 5–7.5 μ long. On cabbage stalks, Batheaston, February 1869."

The foregoing description is quoted from Berkeley and Broome (*loc. cit.*). There do not appear to be any perithecia on the type specimen in Hb. B.M. now. Saccardo (*loc. cit.*) suggested that *N. furfurella* was the same as *N. Keithii*, and Cooke adopted the suggestion in *Grevillea*, XII (1884), 10, but it is doubtful whether either had examined specimens. *Nectria furfurella* has not been reported again.

MELANOSPORA Corda, *Icon. Fung.* I (1837), 24

Perithecia separate, superficial, more or less globose, with a long beak fringed with hyaline setae at the tip, or without a beak, but with a fringe of setae round the ostiolum; asci usually eight-spored, soon

diffuent; ascospores continuous, fuscous, dark brown or black. Wall of perithecium membranous, hyaline or light-coloured, subtransparent, the whole appearing black when the ascospores are mature.

Melanospora caprina (Fr.) Sacc., *Syll. Fung.* II (1883), 462; *Sphaeria caprina* Fr. in *Fl. Danica*, pl. 1859, fig. 2 (1829); *Ceratostoma caprinum* Fr., *Summa Veg. Scand.* (1849), 396; *Melanospora vervecina* (Desm.) Fuckel, *Symb. Mycol.* (1869), 126; *Sphaeria vervecina* Desm. in *Ann. Sci. Nat.* ser. 2, XVII (1842), 103.

Perithecia gregarious, on a rather compact, stout, tomentose, persistent, dark brown or purple-brown subiculum, depressed globose, clothed with a thick, white, woolly layer of tomentum, up to 0.8 mm. diameter (including the tomentum), with a beak, up to 1.8 mm. high, 0.2 mm. diameter below, tapering upwards and terminating in a pencil of hyaline setae; wall of the perithecium yellow, thick, parenchymatous, the exterior cells up to 30μ diameter; wall of the beak yellow, of irregular, vertically elongated cells; hyphae of the tomentum thick-walled or almost solid, about 3μ diameter, variously curled and intertwined, extending from the lower part of the perithecium over the subiculum; asci broadly clavate, $100 \times 20\mu$, spores clustered, or more or less biseriate; ascospores dark brown, broadly oval, or somewhat lemon-shaped, $16-23 \times 9-16\mu$.

On dead wood. Glamis, January 1874 (Stevenson), Hb. Kew.; no locality, December 1873 (Keith), Hb. Kew.; Clunhill, November 1878 (Keith); Forres, on larch, Hb. Kew.; Carlisle, 16 December 1883 (Dr Carlyle), Hb. Kew.; Rudloe, November 1842 (Broome), Hb. Kew. Also recorded from Leigh Woods, Bristol, by Bucknall.

Melanospora chionea (Fr.) Corda, *Icon. Fung.* I (1837), 25, pl. 7, fig. 297; *Sphaeria chionea* Fr., *Syst. Mycol.* II (1823), 446; *Ceratostoma chionea* Fr., *Observ.* II (1818), 340, pl. 7, fig. 2.

Perithecia gregarious, globose or depressed globose, 0.36–0.5 mm. diameter, with a conical beak, 0.36–0.42 mm. high, 0.08–0.1 mm. diameter below, tapering to 0.04 mm. diameter above and terminating in a pencil of hyaline setae; beak and wall of the perithecium yellow or pale brown; beak composed of irregular, vertically elongated cells; wall of the perithecium parenchymatous, woolly with a dense white covering of hyaline, more or less regular, thick-walled hyphae, $2-2.5\mu$ diameter, variously curled and intertwined, in general not extending over the substratum; asci clavate with a long pedicel, sporiferous part, $35-45 \times 14-18\mu$ (Winter), $56 \times 16\mu$ (Fuckel), spores clustered, or more or less biseriate; ascospores oval or subcircular, discoid, dark brown, not apiculate, $10-16 \times 8-12\mu$, $4-6\mu$ thick.

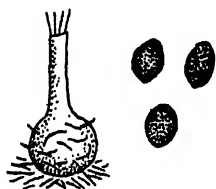


Fig. 5. *Melanospora cirrhata*; perithecium, $\times 60$; ascospores, $\times 300$.

On pine needles. Grantown, November 1878 (Keith), det. Plowright; Orton Moss, April 1881 (Dr Carlyle), Hb. Kew.; Dunkeld, 23 April 1914 (C. Rea), Hb. Kew.; without locality, May 1914 (J. W. Ellis), Hb. Kew. Also recorded for Bulmer, Yorks, in *Fungus Flora Yorks*.

Melanospora Zamiae Corda, *Icon. Fung.* 1 (1837), 24, pl. 7, fig. 297 A; *Ampullaria aurea* A. L. Smith in *Journ. Bot.* xli (1903), 258.

Perithecia superficial, gregarious, globose, 160–400 μ diameter, with a long, cylindrical beak, terminated by a pencil of setae, 170–560 μ high, yellow or reddish, sparsely clothed with white or yellowish flexuose hyphae, wall yellow by transmitted light, or brownish yellow in old specimens; asci broadly clavate, 60–70 \times 28–35 μ ; ascospores lemon-shaped, dark brown to black, 18–23 \times 12–16 μ .

On rotting plants under damp conditions. On dead clover seeds, Norwood, April 1903 (A. L. Smith); on seedlings from seed of Timothy grass infected with *Acremonia atra*, Aberystwith, March 1927 (K. Sampson); on seedlings of Timothy grass, Kew (E. W. Mason); on bananas from the North Riding Laboratory of Pathology and Public Health, October 1931 (E. W. Mason). See Mason, in *Annotated Account of Fungi received at the Imperial Mycological Institute*, List II, fasc. 2 (1933).

Melanospora cirrhata Berk., *British Fungi*, No. 325 (1843); ? *Gibsonia phaeospora* Masee in *Ann. Bot.* xxiii (1909), 366.

Perithecia small, scattered, globose, 0.2 mm. diameter, with a cylindrical, straight or curved beak, 0.25 mm. high, 0.08 mm. diameter, abruptly defined from the perithecium and terminating in a pencil of hyaline setae; wall of the perithecium and beak yellow-brown (in herb. specimens), parenchymatous, membranous, the membrane of irregularly longitudinal cells; perithecium sparsely clothed with pale yellowish spreading hyphae, thick-walled or almost solid, 2.5 μ diameter, which also extend in a circular patch from the base of the perithecium over the substratum; asci not seen; ascospores dark brown, broadly oval, not or slightly apiculate, 13–18 \times 10–13 μ , or circular, 15 μ diameter.

On straw, King's Cliffe, 16 April 1841 (Berkeley), Hb. Kew. and Hb. B.M.

Melanospora leucotricha Corda, *Icon. Fung.* 1 (1837), 25, pl. 7, fig. 297 C.

Perithecia gregarious or scattered, on a cobwebby mycelium which may overrun wide areas, globose, 0.3 mm. diameter, passing into a conoid or subcylindrical beak, 0.35 mm. high, of which the uppermost 0.1 mm. consists of a pencil of hyaline setae; perithecium "translucent yellowish white", yellow-brown in herbarium specimens, wall parenchymatous, stout, about 66 μ thick, covered with loosely intertwined, white, thick-walled hyphae, 2–3 μ diameter, which usually do not spread out radially to the substratum; beak 90 μ diameter below, 75 μ above, thick-walled, yellow-brown, externally glandular with thick-walled, irregularly oval or pyriform

cells, about $12 \times 9 \mu$; asci "oblong, stalked, 4 or 8-spored, sporiferous part $38-50 \mu$ long, $17-23 \mu$ broad" (Winter); ascospores dark brown, broadly ellipsoid, apiculate, $17-23 \times 13-18 \mu$.

"On heaps of rotting leaves, stems, branches, wood, even running over stones."

No British specimens are available. The foregoing description has been drawn up from Rabh.-Winter, no. 2757, Leipzig 1881, on decaying leaves. Recorded on decaying Cherry Laurel leaves, Monaghan (*Trans. Brit. Mycol. Soc.* III, 210); also, on plates exposed over apple orchards, F. M. Carter (*op. cit.* XIX, 146).

Melanospora parasitica Tul., *Sel. Fung. Carp.* III (1865), 10, pl. 3, figs. 11-14.

Perithecia partly immersed in the hyphae of the host, globose, about 0.2 mm. diameter, with a long cylindrical beak, up to 2.5 mm. high, $50-90 \mu$ diameter below, tapering to $20-40 \mu$ above, terminated by a pencil of hyaline setae, at first entirely hyaline, becoming brownish to reddish brown; wall of the perithecium parenchymatous, membranous; wall of the beak composed of somewhat irregular, but more or less parallel hyphae; with stout hyphae arising from the perithecial wall and spreading through the host, and a slight covering of contorted hyphae at the base of the beak; asci clavate, eight-spored, $27 \times 5 \mu$, soon disappearing; ascospores cylindrical, ends truncate, fuscous to black, $5-8 \times 2-2.5 \mu$.

On entomogenous fungi. On *Isaria farinosa*, West Briggs; near King's Lynn, 10 September 1931; Rawcliffe, Yorks, 4 August 1934. On *Beauveria Bassiana*, Dartington, Devon, 25 September 1935. Recorded by Plowright (*Grevillea*, x, 71) on *Isaria farinosa*, North Wootton, September 1880.

Melanospora lagenaria (Pers.) Fuckel, *Symb. Mycol.* (1869), 126; *Sphaeria lagenaria* Pers., *Synopsis* (1801), 58; *Ceratostoma lagenarium* (Pers.) Fr., *Observ.* II (1818), 341; *Melanospora lagenaria* var. *tetraspora* Rehm in *Hedwigia* (1891), 259.

Perithecia scattered or clustered, sometimes connate, globose, or conoid above and merging gradually into the beak, yellowish brown, becoming dark brown, 0.36-0.5 mm. diameter, with a beak up to 1.5 mm. high, 0.2 mm. diameter below, tapering to 0.1 mm. diameter above, straight or curved, terminating in a pencil of hyaline setae; wall of the perithecium stout, subopaque, hispid above with short yellowish hairs, clothed below with longer hyphae which sometimes envelop adjacent perithecia; beak glabrous, dark red-brown in old specimens; asci stalked, broadly clavate, sporiferous part $36 \times 13 \mu$, eight-, four-, or three-spored in the same perithecium, spores clustered; ascospores dark brown, ellipsoid, sometimes inequilateral, slightly apiculate, $13-22 \times 7-14 \mu$.

On old *Polypori*. On *Polyporus adustus*, Queen's Cottage, Kew, April 1888 (Cooke), Hb. Kew.; on *Polyporus*, Epping Forest, 19 October 1918 (Wakefield), Hb. Kew. Also recorded on decaying ? *Stereum*, Bushey Park (*Trans. Brit. Mycol. Soc.* II, 93), and on *Polystictus versicolor*, Eglinton, Ayrshire (*op. cit.* VI, 47).

Melanospora brevirostris (Fuckel) v. Höhnelt in *Sitzb. k. Akad. d. Wiss. Wien*, CXXII (1914), 94; *Ceratostoma brevirostre* Fuckel in *Bot. Zeit.* XIX (1861), 250, pl. 10, fig. 4; *Melanospora Zobelii* Fuckel non Corda, *Symb. Mycol.* (1869), 127; *Ceratostoma Helvellae* Cooke in *Grevillea*, I (1873), 175; *Melanospora Helvellae* (Cooke) Sacc. in *Michelia*, I (1878), 283.

Perithecia at first ovoid and astomate, becoming subglobose with a conical apex which opens by a pore surrounded by hyaline setae, which sometimes develop into a cylindrical, hyaline beak, up to 180μ high; perithecia superficial, scattered or crowded, up to 0.36 mm. diameter, wall parenchymatous, stout, brown, smooth, or with a few adherent brown hyphae; asci not seen; ascospores oval, sometimes acuminate, sometimes inequilateral, ends truncate, dark brown, $22-31 \times 13-16\mu$, with some globose, $13-16\mu$ diameter.

On the disc of *Sepultaria arenosa* and allied Discomycetes, sometimes entirely covering it. On *Peziza sepulta*, without locality, November 1858 (Rev. H. Higgins), Hb. Kew. ex Hb. Currey; on *Peziza hemisphaerica*, Eastbourne, February 1873 (C. J. Muller), Hb. Kew.; Sandhills, South Lancashire, June 1920 (W. G. Travis), Hb. B.M. Also recorded, as *M. Zobelii*, on *Sepultaria arenicola* (Lév.) Mass., Wallasey Sandhills, 25 November 1913 (J. W. Ellis), in *Trans. Brit. Mycol. Soc.* IV, 314.

Melanospora damnosa (Sacc.) Lindau in Engler-Prantl, *Naturl. Pflanzenf.*, Teil I, abt. I, 353 (1897); *Sphaeroderma damnosum* Sacc., in Berlese, *Rivista Pat. Veget.* (1895), 9, pls. 7 and 8.

Perithecia concealed beneath the raised periderm, only the white pencil of convergent setae visible externally. Perithecia globose, about 250μ diameter, ochraceous or reddish, with a short cylindrical beak about 50μ high, crowned with a white conical pencil of hyaline setae, about 350μ high; wall yellow or brownish yellow by transmitted light, clothed with hyaline or yellowish hyphae; asci broadly clavate, four-spored, about $45 \times 18\mu$; ascospores broadly ellipsoid or subglobose, not or slightly apiculate, dark brown, black in mass, $16-21 \times 12-18\mu$.

On elm twigs, Kew, kept in a damp chamber, July 1932 (E. W. Mason).

Melanospora fimbriata (Rostrup) Petch, comb. nov.; *Sphaeroderma fimbriatum* Rostrup, *Oest. Groenl. Svampe* (1894), 25.

Subiculum obsolete; perithecia very small, globose, reddish, ostiolum fimbriate; asci cylindraceo-clavate, $100-110 \times 20\mu$; ascospores ellipsoid, fuscous, $20 \times 11-12\mu$ (Rostrup).

On dung of guinea-pig, Kew (Massee & Salmon). No specimens available.

Melanospora sphaerodermoides Grove in *Journ. Bot.* XXIII (1885), 132, pl. 256, fig. 4.

Perithecia superficial, scattered, almost smooth, pale yellow-brown, then brown, subglobose, conoid above, about 0.35 mm. diameter, 0.4 mm. high, opening by a pore surrounded by a short collar, with

hyaline setae at the upper edge of the collar; setae $50-60\mu$ long, rigid, acute, continuous, up to 5μ diameter below, tapering to the tip; wall parenchymatous, rigid, of large hexagonal cells; asci obovate-clavate, stalked, eight-spored, $80-90 \times 30\mu$; ascospores oval or inequilaterally oval, or subcymbiform, ends truncate, with a germ pore, sometimes shortly acuminate, black, $27-34 \times 13-16\mu$.

On culms of *Heracleum*, Warwickshire (Grove); on decaying stalks of *Brassica*, North Wootton, November 1935.

Melanospora Zobelii (Corda) Fuckel, *Symb. Mycol.* (1869), 127; *Microthecium Zobelii* Corda, *Icon. Fung.* v (1842), 74, pl. 8, fig. 53; *Sphaeria (Hypocrea) Zobelii* (Cda.) Tul., *Fungi Hypogaei* (1851), 186, pl. 13, fig. 1; *Ceratostoma Zobelii* (Cda.) Berk., *Outlines* (1860), 402.

Perithecia membranous, immersed or partly immersed, at first globose and astomate, becoming globose with a conical apex and opening by an apical pore; setae not observed; ascospores almost black, lemon-shaped, ends truncate, $21 \times 12\mu$ (Corda), $23 \times 16\mu$ (Tulasne).

On or in the hymenium of truffles. Recorded for Britain by Berkeley (*loc. cit. supra*) without locality; not represented in British Herbaria. When specimens are available, this species may prove to be identical with *Melanospora brevirostris*, as was maintained by Fuckel.

SPHAERODERMA Fuckel, *Symb. Mycol. App.* III (1875), 22

Perithecia and ascospores as in *Melanospora*, but without a beak or setae round the ostiolum.

Sphaeroderma fusisporum Petch in *Naturalist*, (1936), 58.

Perithecia scattered, superficial, orange, globose, up to 0.3 mm. diameter, at first astomate, but developing a papillate ostiolum and subsequently a collar round the orifice, glabrous, or with a few adpressed hyphae; wall yellow by transmitted light, parenchymatous, of large cells; asci eight-spored, clavate, $66 \times 12\mu$, soon diffuent, spores obliquely uniseriate, or biseriate; ascospores lanceolate, fuliginous, apices obtuse, or truncate with a germ pore, continuous, with a large central gutta, $20-24 \times 6-9\mu$. Perithecia appearing black from the apex downwards as the spores mature.



Fig. 6. *Sphaeroderma fusisporum*; perithecium, $\times 33$; ascospore, $\times 400$.

On *Isaria farinosa*, Saltaire, Yorks, September 1935; North Wootton, Norfolk, October 1936.

Sphaeroderma episphaeria (Phil. & Plowr.) Sacc., *Syll. Fung.* II (1883), 560; *Melanospora episphaeria* Phil. & Plowr. in *Grevillea*, x (1881), 71, pl. 158, fig. 2; *Microthecium episphaerium* (Phil. & Plowr.) v. Höhnelt in *Sitzb. k. Akad. d. Wiss. Wien*, CXXXIII (1914), Abt. 1, 98;

Sphaerodes episphaerium (Phil. & Plowr.) Clements, *Genera of Fungi* (1909), 44 and 173; *Vittadinula episphaeria* (Phil. & Plowr.) Sacc., in Shear and Clements, *Genera of Fungi* (1931); *Sphaeroderma epimyces* v. Höhnelt, *Fragmente zur Mykologie*, no. 105 (1907).

Perithecia superficial, globose, 0.25–0.35 mm. diameter, becoming globose with a papillate apex and opening by an apical pore, at first hyaline, then appearing black; wall in dried specimens brownish yellow; asci cylindrical, then pyriform, soon disappearing; spore-cluster ovate, about $66 \times 22 \mu$; ascospores lemon-shaped, ends truncate, at first hyaline with vacuolate contents, then almost black and reticulated with a wide-meshed network of narrow lines, $25\text{--}34 \times 12\text{--}18 \mu$.

On *Hypomyces ochraceus* (Pers.) Tul. North Wootton and Holt House Wood, near King's Lynn, October 1880 (Plowright), Hb. Kew.; Ashwicken, 28 October 1896 (Plowright).

Sphaeroderma Hulseboschii Oud., *Contr. Flor. Pays Bas* xi (1886), 23.

Perithecia superficial, without any subiculum, subglobose, 0.7 mm. diameter, pale ochraceous, ostiolum short, obtusely conical; asci pyriform, $50 \times 25 \mu$, eight-spored; ascospores lemon-shaped, subolivaceous, $19\text{--}21 \times 11\text{--}12 \mu$ (Oudemans).

On rabbit dung, Leith Hill, Surrey (Massee & Salmon). No specimens available. Probably not different from *Sphaeroderma fimicolum* (E. C. Hans.) Sacc.

NECTRIA Fries, *Summa Veg. Scand.* (1849), 387 (in part)

Perithecia superficial, separate, but caespitose on a superficial or erumpent, parenchymatous, bright-coloured stroma, bright-coloured, soft; asci usually eight-spored, sometimes polysporous by budding from the ascospores; ascospores hyaline, one-septate. In some species, scattered perithecia may occur in company with the normal stromatic clusters.

Nectria cinnabarina (Tode) Fr., *Summa Veg. Scand.* (1849), 388; *Sphaeria ochracea* Grev. & Fr., *Elenchus*, II (1828), 79; *Nectria ochracea* (Grev. & Fr.) Fr., *Summa Veg. Scand.* (1849), 387; *Nectria fuscopurpurea* Wakefield in *Kew Bulletin* (1918), 232.

Perithecia crowded on an erumpent pulvinate stroma, cinnabar or purple-red, darkening with age, globose, up to 0.4 mm. diameter, covered with coarse warts, ostiolum naked, papillate; asci cylindrico-clavate, eight-spored, spores biserial; ascospores oblong or narrow-oval, often inequilateral, ends rounded, one-septate, $12\text{--}25 \times 4\text{--}9 \mu$; paraphyses stout, septate, branched.

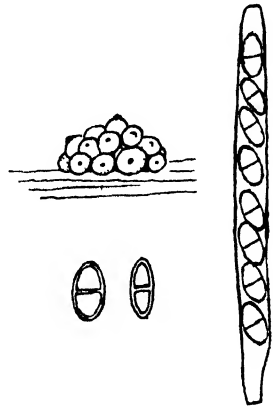


Fig. 7. *Nectria coccinea*; group of perithecia, $\times 10$; ascus, $\times 500$; ascospores, $\times 500$.

Conidial stage, *Tubercularia vulgaris* Tode, pulvinate, red, sub-translucent when moist, pink when dry; conidia oblong-oval or sub-cylindric, ends rounded, straight or slightly curved, $6-9 \times 2-2.5 \mu$.

Common on dead branches.

In the type of *Nectria fuscopurpurea*, the perithecia are almost smooth, turbinate and concave or strongly umbilicate when dry, and some of the ascospores are up to 33μ long, and two to three septate. The majority of the ascospores, however, are those of *N. cinnabarina*, and it would appear to be an abnormality of that species. There is a similar specimen in Hb. Kew., ex Hb. Berkeley, without date or locality, labelled "*Sphaeria cucurbitula* Tode, 47", by Berkeley and I have recent specimens from Norfolk and Yorkshire.

Nectria Ralfsii B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, XIII (1854), 467.

Perithecia in small groups of about six on a small superficial stroma which is yellow internally, depressed globose, 0.33 mm. diameter, orange, covered with large, confluent, paler warts composed of more or less globose, thick-walled cells, $9-12 \mu$ diameter, naked round the minute, conical ostium; asci clavate, $70 \times 14-16 \mu$, spores biserial; ascospores oval, narrow-oval, oblong-oval, or fusoid, hyaline, one-septate, the lower cell often narrower and tapering, $16-27 \times 6-9 \mu$, with some, one-septate, broadly ellipsoid, $12-14 \times 9 \mu$, and some globose, one-celled, $9-12 \mu$ diameter.

On ? beech, Penzance (Ralfs), Hb. Kew. and Hb. B.M.; on gorse, Penzance (Ralfs), Hb. Kew. and Hb. B.M.; Southampton, February 1910 (Rayner), Hb. B.M.; on *Acer*, Berry Pomeroy, Devon, September 1935 (E. M. Wakefield) Hb. Kew. Also recorded from Goole (*Naturalist*, September 1881).

Nectria subquaternata B. & Br. in *Journ. Linn. Soc.* XIV (1873), 116; *Nectria squamuligera* Sacc., *Fungi Veneti*, ser. 4, 22 (1875); *Nectria Keithii* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 4, XVII (1876), 144; *Nectriella Keithii* (B. & Br.) Sacc. in *Michelia*, I (1878), 278.

Perithecia clustered on a thin, fleshy, yellow or yellow-brown stroma, or scattered, globose, 0.25 mm. diameter, pale yellow-brown to dark brown, subtranslucent when moist, at first with a conspicuous ring of white parenchymatous warts round the ostium, the warts subsequently becoming scattered and of the same colour as the perithecium; ostium minute, conical, naked; wall stout, of small cells, yellow-brown; asci subcylindric or clavate, almost sessile, apex truncate, $48-60 \times 7-10 \mu$; paraphyses filiform, branched; ascospores narrow-oval or subcylindrical, sometimes inequilateral, one-septate, ends obtuse, $9-16 \times 3-5 \mu$.

On decaying stalks of *Brassica*. Batheaston, 5 March 1877 (Broome), Hb. B.M.; Forbes, (Keith), Hb. Kew.; North Wootton, King's Lynn, October 1935.

The specimens on stalks of *Brassica* differ in colour from Ceylon and Italian specimens, the latter being pale flesh-coloured when fresh, and yellow or pallid when dry. A specimen in Hb. B.M., labelled "*N. squamuligera*, Mulgrave Woods, July 1910, T. Gibbs", is *Nectria Peziza* (Tode) Fr.

Nectria inventa Pethybridge in *Trans. Brit. Mycol. Soc.* vi (1920), 107.

Perithecia gregarious, on a small, dark brown, superficial stroma, which is sometimes situated in a decaying patch of *Acrostalagmus cinnabarinus*, globose, 300–500 μ diameter, blackish brown, bearing short, rigid, septate hairs on the upper half; hairs hyaline, up to 175 μ high, 6 μ diameter below, tapering upwards, septate, regular, or sometimes with an oval swelling at varying heights, apex acute, sometimes lanceolate; wall of the perithecium of small cells, yellow-brown by transmitted light; asci cylindrical or cylindrico-clavate, 60–100 \times 4–6 μ , eight-spored; paraphyses filiform, 150 \times 3–4 μ , diffluent; ascospores obliquely uniseriate, oblong, ends rounded, hyaline, one-septate, 8–12 \times 3–5 μ .

Conidial stage, *Acrostalagmus cinnabarinus* Corda.

On rotting potatoes, Ireland (Pethybridge). On decaying stalks of *Brassica*, North Wootton, King's Lynn, November 1935, October 1936.

Nectria coccinea (Pers.) Fr., *Summa Veg. Scand.* (1849), 388.

Perithecia usually clustered on a red stroma, which is yellow internally, sometimes scattered without a stroma, globose, 0.25–0.3 mm. diameter, with an evident conical ostium, yellowish red, ostium darker, becoming dark red, smooth; perithecial wall of rather large cells; asci cylindrical or narrow-clavate, apex truncate and thickened, with a central pore, spores uniseriate, or obliquely uniseriate, or biseriate above, the uppermost spore sometimes some distance below the apex, asci sometimes clavate with a rounded apex and the uppermost spore in contact, but the original thickened apex persisting as a small disc, 66–95 \times 8–12 μ ; paraphyses surrounding the mass of asci, broad, septate, and branched as in *Dialonectria galligena*, but thin-walled and soon diffluent; ascospores narrow-oval, oblong-oval, or sub-fusoid, ends rounded, one-septate, sometimes slightly constricted, hyaline, minutely warted, 12–19 \times 5–7 μ .

On dead branches; common.

Nectria punicea (K. & Schm.) Fr., *Summa Veg. Scand.* (1849), 387; *Nectria ditissima* Tul., *Sel. Fung. Carp.* III (1865), 72.

Perithecia caespitose on a red stroma which is subgelatinous when moist, globose or globoso-conoid, sometimes distinctly conoid when effete, up to 0.36 mm. high, 0.3 mm. diameter, with a minute, papillate, scarcely elevated ostium, bright red, becoming purple-red, dark red and subtranslucent when moist, pruinose, becoming dull brownish red and smooth, sometimes brownish yellow in old specimens; asci at first subfusoid, then clavate, apex at first obtuse or subtruncate, not thickened, finally rounded, 90–100 \times 10–12 μ , spores obliquely uniseriate, becoming biseriate above; paraphyses broad,

septate and branched as in *Dialonectria galligena*, but thin-walled and soon diffluent; ascospores narrow-oval, or oblong-oval, sometimes inequilateral, one-septate, hyaline, minutely warted, $12-20 \times 4-7 \mu$.

Conidial stage; the conidial fungus which occurs commonly with this species is *Tubercularia minor* Link, Sp. Plant., vi, 2 (1825), 100, *T. crassostipitata* Fuckel, *Symb. Mycol.* (1869), 180, but there is some doubt whether it is the conidial stage of *N. punicea*.

On *Rhamnus frangula*; Highgate, Cooke, *Fung. Brit. Exsicc.* no. 370, and ed. II, no. 475; Bishop's Wood, February 1866 (Cooke), Hb. Kew. On laburnum, Leicester (F. T. Mott) and Kingscliffe (Berkeley), Hb. Kew., det. Ehrlich. On ash, Forden, Vize, *Microf. Brit.* no. 152, det. Ehrlich. On holly, Forden, Vize, *Microf. Brit.* no. 373, and Shere (Cooke), Hb. Kew, det. Ehrlich., etc. On beech, North Wootton, March 1937. On broom, North Wootton, November 1933; Norwich, October 1934. On ivy, Wheatfen Broad, Norfolk, January 1936 (E. A. Ellis).

Nectria sinopica Fr., *Summa Veg. Scand.* (1849), 388.

Perithecia crowded on a pale red erumpent stroma (yellow internally), globose, 0.25–0.3 mm. diameter, with a small conical ostiolum, becoming umbilicate when old, at first light red or brick-red, subtranslucent, pruinose with yellow granules, ostiolum darker, becoming dark red, and finally purple-red or blackish; asci cylindrical, $60-90 \times 6-9 \mu$; ascospores ellipsoid or oblong-oval, sometimes inequilateral, $9-15 \times 4-6 \mu$, hyaline, becoming pale yellow.

Pycnidial stage, *Zythiostroma Mougeotii* (Fr.) v. Höhn., pycnidia subcortical, ostiolum erumpent, or exposed by abscission of the cortex, scattered, red, soft, subtranslucent, subglobose or conoid, 0.25–0.4 mm. diameter, ostiolum cylindrical, dark red; pycnosporos oblong-oval, $2.5-3.5 \times 1 \mu$. With the perithecial stage.

On dead stems of ivy; generally distributed.

Nectria cucurbitula Sacc. in *Michelia*, I (1878), 409; *Sphaeria cucurbitula* var. *β. nigrescens* Tode, *Fung. Meckl.* II (1791), 39; *Nectria cucurbitula* (Tode) Fr. (in part), *Summa Veg. Scand.* (1849), 388.

Perithecia caespitose on an immersed or feebly erumpent stroma, at first brick-red, then blood-red, later becoming purple brown and finally blackish, globose, with a small conical ostiolum, smooth, generally shining, about 0.3 mm. diameter; asci cylindrical or slightly clavate, subsessile, $90-110 \times 7-9 \mu$, apex truncate and thickened, spores uniseriate or obliquely uniseriate; ascospores oval or oblong-oval, ends rounded or sometimes somewhat triangular, hyaline, one-septate, not or slightly constricted, $12-15 \times 5-6 \mu$.

On coniferous logs. On *Picea excelsa*, near Braco, Perthshire, 31 March 1935; on *Pinus sylvestris* and ? *Larix europaea*, Benmore Forestry Reserve, Argyllshire, 30 March 1935 (C. G. C. Chesters), det. Ehrlich.

This species has been confused with *Nectria Coryli*, and most of the British records refer to the latter.

[*Nectria Solani* Reinke & Berth., *Zersetz. d. Kartoffel* (1879), 39.

Perithecia crowded on an erumpent convoluted stroma, globosconoid, with a conical ostiolum, pale ochraceous, almost white, or orange-red; asci cylindrico-clavate or broadly clavate, $45-65 \times 7-11 \mu$; ascospores oblong or oblong-oval, ends acute, hyaline, one-septate, constricted at the septum, $8-14 \times 4-6 \mu$.

Conidial stage. Reinke and Berthold state that the conidial stage of *Nectria Solani* is *Spicaria Solani* de Bary. Their figure shows a *Gliocladium*, with narrow-oval or lanceolate conidia, $4-5 \times 3 \mu$.

Pethybridge (*Trans. Brit. Mycol. Soc.* vi (1920), 105) states that there is no reliable record of the occurrence of *Nectria Solani* in the British Isles.]

Nectria Coryli Fuckel, *Symb. Mycol.* (1869), 180; *Nectria cucurbitula* Currey in *Trans. Linn. Soc.* xxii (1858), 282, pl. 49, fig. 178; *Chilonectria cucurbitula* (Curr.) Sacc. (in part) in *Michelia*, i (1878), 280.

Perithecia erumpent, caespitose, orange, yellowish red, or red, sometimes covered with yellow particles, becoming dark red, blackening when old, globose, 0.3 mm. diameter, with a minute conical ostiolum, darker than the rest of the perithecium, becoming cup-shaped or turbinate when dry; asci narrow-clavate, apex truncate and thickened, $75-95 \times 5-7 \mu$, spores uniseriate or obliquely uniseriate, or partly biseriate; paraphyses linear, diffuent; ascospores narrow-oval or subcymbiform, hyaline, one-septate, not or slightly constricted, $10-18 \times 4-5 \mu$, budding in the ascus and producing myriads of minute, curved, allantoid hyaline sporidia, $2-4 \times 0.75-1 \mu$; the perithecia may also contain subcylindric, three-septate, hyaline spores, $40-42 \times 5-7 \mu$.

On ash (Berkeley), Hb. Kew. On privet, Lucknam, March 1850 (Broome), Hb. B.M. On hazel, Batheaston, February 1859 (Broome), Hb. B.M.; Rokeby, September 1933. On alder, Booton Common, Norfolk, March 1936 (E. A. Ellis). Also recorded on *Ulex*, Leziate Fen, King's Lynn (B.M.S.).

Nectria Aquifolii (Fr.) Berk., *Outlines*, etc. (1860), 393; *Sphaeria Aquifolii* Fries, *Elenchus*, ii (1828), 82; *Nectria inaurata* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, xiii (1854), 467; *Aponectria inaurata* (B. & Br.) Sacc. in *Michelia*, i (1878), 296.

Perithecia erumpent, caespitose on an immersed or erumpent stroma which is yellow or brownish internally, pale red becoming dark red, sometimes naked, sometimes strongly encrusted with yellow or greenish yellow granules, blackening when old, globose, 0.25-0.4 mm. diameter, with a dark brown or almost black papilla, sometimes becoming flat-topped, broadly umbilicate and subturbinate when dry; asci cylindrical or clavate, $60-85 \times 7-11 \mu$, eight-spored, or containing numerous sporidia; ascospores oval, one-septate, $9-15 \times 4-7 \mu$; sporidia cylindric or narrow-oval, straight or curved, $4-6 \times 1-1.5 \mu$.

On dead branches and stems of holly; generally distributed, but often recorded in error for *N. punicea* and *N. coccinea*, both of which occur on holly.

Nectria mammoidea Phil. & Plowr. in *Grevillea*, III (1875), 126.

Perithecia caespitose, frequently scattered, red or yellowish red, with a dark red (appearing black) apical area, becoming entirely dark red or purple-brown when old, sometimes pruinose or furfuraceous, globose, up to 0.5 mm. diameter, smooth, not collapsing, ostiolum minute, conical; wall rigid, tough, with a parchment-like layer which often persists after the decay of the rest of the perithecium; asci cylindrical or narrow-clavate, almost sessile, 90–130 × 10–14 μ , spores uniseriate or obliquely uniseriate; paraphyses linear; ascospores narrow-oval or subcymbiform, sometimes broadly fusoid, ends rounded, one-septate, hyaline becoming pale yellow, very minutely warted, 14–25 × 6–9 μ .

Var. *Rubi* Weese in *Zeitschr. Garungsphys.* I (1912), 128; *Nectria Rubi* Osterw. in *Ber. deutsch. Bot. Gesell.* XXIX (1911), 620, tab. xxii.

Perithecia as in the type, but spores smaller, 15.9–18.6 × 4.6–5.2 μ (Osterwalder).

On gorse; also on elm, etc. North Wootton, November 1873, 1874, Plowright, *Sphaer. Brit.* II, 5; Ercall, January 1874 (Phillips), Hb. B.M.; Scarborough (Massee) Hb. B.M.; Mulgrave, May 1911, Hb. B.M.; Rokeby, September 1933; Matlock Bath, June 1935; Dartington, September 1935. Also recorded from Walton Hill and Leigh Wood, July 1881 (Bucknall); Tintern (B.M.S.); Leziate (B.M.S.).

Var. *Rubi*. On roots of bramble, Batheaston, April 1870 (Broome), Hb. B.M.; North Wootton, February 1937. On raspberry, recorded for Scotland by Alcock (*Trans. Bot. Soc. Edinb.* XXIX (1925), 197), for Ireland by Pethybridge (*Trans. Brit. Mycol. Soc.* XII (1927), 20), and for Worcestershire by Nattrass (*tom. cit.*, p. 23).

Nectria Magnusiana Rehm, *Ascomyceten*, no. 436.

Perithecia caespitose on an inconspicuous stroma, globose, 0.25 mm. diameter, slightly papillate when young, collapsing and becoming regularly cup-shaped, bright red, becoming dull red, and then blackening, minutely rugose; asci cylindrical, almost sessile, apex rounded and thickened, 75–90 × 7–8 μ , spores uniseriate or obliquely uniseriate; ascospores ellipsoid, ends rounded, smooth, one-septate, hyaline becoming reddish, 9–14 × 5–7 μ .

Conidial stage, *Dendrodochium epistroma* v. Höhn., *Fragm. z. Mykol.* VI, no. 284. Sporodochia minute, pulvinate, or confluent in patches, pale red when fresh, orange-red or blood-red when dry, rather soft; conidiophores branched above, conidia terminal; conidia hyaline, cylindrical, straight or curved, 3–7 × 0.75–1 μ .

On *Diatrypella* on birch, with the conidial stage, Logan Woods, near Sandhead, Wigtownshire, August 1936 (C. G. C. Chesters); Becca Park, Aberford, Yorks. October 1937 (W. G. Bramley); conidial stage only, Great Bear Park, Warwickshire (Grove); North Wootton, near King's Lynn, November 1935, February 1936, etc.

Nectria ochroleuca (Schw.) Berk. in *Grevillea*, iv (1875), 16; *Sphaeria ochroleuca* Schw. in *Trans. Amer. Phil. Soc. n.s.* iv (1832), 204.

Perithecia caespitose, on a small, inconspicuous, erumpent, pulvinate stroma, which is sometimes floccose with white conidiophores, globose, small, about 0.2 mm. diameter, minutely rugose, pale yellow or almost white, opaque, dark round the minute, conical ostiolum; asci sessile, narrow-clavate or subcylindrical, apex truncate and impressed, spores uniseriate, or sometimes biseriate above, $55-66 \times 5-6 \mu$; paraphyses not seen; ascospores narrow-oval, often inequilateral, or fusoid, straight or slightly curved, $10-14 \times 3-4 \mu$, sometimes $15-18 \times 3.5 \mu$ and two- or three-septate.

Conidiophore verticillate (*Verticillium tubercularioides* Speg.); conidia elliptical, hyaline, $5-8 \times 3 \mu$ (Seaver).

Coed Coch, October 1880 (Cooke); on wych elm, Forden, Vize no. 2, Hb. Cooke; Baslow, September 1919 (E. M. Wakefield); all in Hb. Kew., det. or confirmed Ehrlich.

Nectria pallidula Cooke in *Grevillea*, xvii (1888), 3.

Perithecia scattered or crowded, on an erumpent, pulvinate, subgelatinous stroma which shrinks and blackens on drying, small, subglobose with a minute, conical ostiolum, or ovoid, 0.18-0.25 mm. diameter, sordid ochraceous, subtranslucent when fresh, becoming dull ochraceous, or pale yellow and opaque when dry, faintly rugose, ostiolum situated on a darker, subtranslucent area on the yellow, opaque examples; stroma yellowish internally, fibroso-gelatinous, base of the perithecium sometimes embedded in the stroma; cells of the perithecial wall filled with a yellow oil when fresh; asci almost sessile, clavate, $50-60 \times 7-9 \mu$, apex truncate and impressed; spores obliquely uniseriate, becoming biseriate above; paraphyses linear; ascospores narrow-oval, oblong-oval, or subfusoid, ends rounded, one-septate, hyaline, $9-14 \times 3-4 \mu$.

On beech, Carlisle (Cooke), Hb. Kew.; on broom, North Wootton, 22 October 1934.

On the available specimens, *N. pallidula* differs from *N. ochroleuca* in its subgelatinous stroma, its shorter and broader asci, and in the presence of oil globules in the perithecial wall. Its perithecia vary from dull ochraceous and subtranslucent to pale yellow and opaque, the latter resembling exactly those of *N. ochroleuca*. It is possible that a wider range of specimens may prove them identical.

Nectria citrino-aurantia (de Lacr.) Desm., *Pl. Crypt. France* (1860), no. 778; *Calonectria citrino-aurantia* (de Lacr.) Sacc. in *Michelia*, i (1878), 314.

Perithecia small, crowded on an erumpent, pulvinate, pale yellow, subgelatinous stroma, often with the base immersed, oval or conoid, about 0.1 mm. diameter, 0.15 mm. high, pale yellow or ochraceous, subtranslucent, umbilicate when dry, smooth; asci clavate, $30-40 \times 5 \mu$; ascospores cylindrical, subfusoid, or narrow-oval, hyaline, one-septate, $6-9 \times 2-2.5 \mu$, with a few globose, continuous, 3μ diameter.

On willow twigs, Batheaston, December 1873 (Broome), Hb. Kew. and Hb. B.M.

SPHAEROSTILBE Tulasne, *Sel. Fung. Carp.* I (1861), 130

Perithecia and ascospores as in *Nectria*, but usually accompanied by an erect *Stilbella* or *Microcera* conidial stage; if the conidial stage is absent, the perithecia appear as *Nectria* (in *S. aurantiaca*), or as *Hyphonectria* (in *S. flammea*).

Sphaerostilbe aurantiaca Tul., *Sel. Fung. Carp.* I (1861), 130, and III (1865), 101.

Perithecia arising from the base of the conidial form, or independently, caespitose, globose, up to 0.4 mm. diameter, slightly umbilicate, pruinose or minutely fibrillose, yellowish red becoming purple-red; asci broadly clavate, with a long, thin, or a short, stout pedicel, $90-130 \times 18-22 \mu$; ascospores oval, one-septate, not or slightly constricted, $20-29 \times 8-10 \mu$.

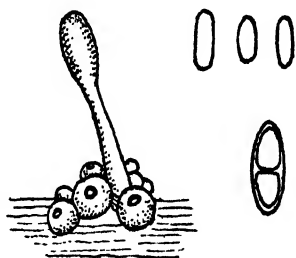


Fig. 8. *Sphaerostilbe aurantiaca*; perithecia and *Stilbella*, $\times 12$; conidia, $\times 400$; ascospore, $\times 400$.

Conidial stage, *Stilbella aurantiaca* (Bab.) Lindau in *Rabh. Krypt-Flora* IX (1908), 298; *Stilbum aurantiacum* Babington, in Berkeley in *Ann. Mag. Nat. Hist.* VI (1841), 432, pl. 12, fig. 14. Solitary or fasciculate, up to 2 mm. high, rarely terete, usually compressed and laterally expanded; stalk red-brown, becoming paler upwards, thickened at the base, smooth, subtranslucent; head subglobose or clavate, red, conidiophores branched, branches long, slender, $1.5-2 \mu$ diameter, conidia terminal; conidia ellipsoid or oblong-oval, ends rounded, sometimes slightly attenuated below and subapiculate, continuous, hyaline, $14-18 \times 6-8 \mu$. The stalk may be lacking, and the conidial fructification then takes a *Tubercularia* form.

On elm. Perithecial stage, Elmhurst, 28 December 1852 (Berkeley); Weybridge, August 1856 (Currey); Salisbury, October 1866 (Cooke); Oundle, July 1868 (Berkeley); all in Hb. Kew., with a duplicate of the first in Hb. B.M. Also recorded from Brandon, 14 October 1881 (Plowright), and from Long Ashton (B.M.S.).

Sphaerostilbe flammea Tul., *Sel. Fung. Carp.* I (1861), 130.

Perithecia usually clustered on a well-developed plectenchymatous stroma, bright orange-red, darker round the ostiolum, globose, 0.3 mm. diameter, glabrous, slightly rugose, opaque, usually collapsing centrally, ostiolum minute, conical; asci cylindrical, almost sessile, eight-spored, $90-110 \times 8-10 \mu$, spores obliquely uniseriate; ascospores ellipsoid, obtuse, one-septate, hyaline or yellowish, minutely warted, $12-19 \times 5-8 \mu$.

Conidial stage, *Microcera coccophila* Desm. in *Ann. Sci. Nat.* ser. 3, x (1848), 539; *Atractium flammeum* Berk. & Rav. in *Ann. Mag. Nat.*

Hist. ser. 2, XIII (1854), 461. Synnemata arising from a white or pinkish stroma round the scale, stilboid, up to 2.5 mm. high, more usually clavate or conical, up to 0.6 mm. high, 0.25 mm. diameter, or flattened pulvinate, up to 0.75 mm. long, 0.5 mm. broad, orange-red to blood-red, usually clothed with erect fascicles of hyphae at the base; conidia fusiform, straight, or straight with falcate tips, or slightly curved, up to eleven-septate, $50-105 \times 5-7 \mu$.

On scale insects, usually (in the British Isles) on *Chionaspis salicis* on ash and willow. Perithecial stage, Trengwainton, Penzance, 14 December 1869, Hb. B.M. Conidial stage, Penzance, Hb. Kew. and Hb. B.M.; near Ryde, Isle of Wight, Hb. B.M.

Sphaerostilbe flavo-viridis Fuckel, *Symb. Mycol. Nachtr.* 1 (1871), 310.

Perithecia superficial, globose with a papillate or subcylindrical apex, 0.27–0.36 mm. high, 0.18–0.3 mm. diameter, yellowish red, then blood-red, finally becoming dark red, apex darker, minutely rugose, glabrous at the apex, rarely collapsing, scattered or gregarious, sometimes arising from a yellow-green mass, sometimes grouped at the base of the yellow-green stilboid conidial stage; asci subcylindrical, $90 \times 6-8 \mu$, or clavate, $70-80 \times 9-12 \mu$, apex at first truncate, spores uniseriate or obliquely uniseriate; paraphyses linear, diffuent; ascospores ovate, or oval, or oblong-oval, ends rounded, one-septate, hyaline, becoming pale brown and minutely warted, $9-14 \times 5-7 \mu$, with a few globose, not septate, $6-10 \mu$ diameter.

Conidial stage, *Atractium flavo-viride* Sacc. Stalk generally simple, acicular, 280μ high, sparsely floccose at the base, yellowish green, apex paler, with a globose white head; conidia continuous or one-septate, hyaline, oblongo-fusiform, or subclavate, straight or curved, $8.5-17 \times 3 \mu$, or fusiform, curved, three- to seven-septate, $32-50 \times 4-4.5 \mu$. According to Wollenweber, this is *Fusarium melanochlorum* (Casp.) Sacc.

Portbury, January 1845 (Broome), Hb. B.M.; on ash, with *Diplodia* sp., Woodchester, 13 May 1934 (E. W. Mason), Hb. Kew. det. Wollenweber; on elm logs, North Wootton, 1 February 1935; on larch, Worlingham Wood, East Suffolk, April 1936 (E. A. Ellis): Allerthorpe Common, Yorks, September 1937 (W. G. Bramley). Usually associated with a Pyrenomycete, often *Melanomma Pulvispyrius* (Pers.) Fuckel.

DIALONECTRIA Cooke in *Grevillea*, XII (1884), 77 and 109

Perithecia superficial, separate, smooth, without a stroma or a continuous subiculum, bright-coloured; asci eight-spored; spores one-septate, hyaline.

Dialonectria Peziza (Tode) Cooke in *Grevillea*, XII (1884), 110; *Sphaeria Peziza* Tode, *Fung. Meckl.* II (1791), 46; *Nectria Peziza* (Tode) Fr., *Summa Veg. Scand.* (1849), 388; *Nectria aurea* Cooke non Grev. in *Grevillea*, VIII (1879), 9; *Nectria epigaea* Cooke in *Grevillea*, VIII (1879), 10.

Perithecia superficial, scattered or gregarious, at first globose, then usually collapsing and becoming cup-shaped, 0.25–0.4 mm. diameter, ochre-yellow, then orange or red, at first with pale yellow or white hyphae arising from the sides and base of the perithecium and spreading radially over the substratum, but usually soon becoming naked; asci cylindrico-clavate, $50-90 \times 6-10 \mu$; ascospores broadly ellipsoid, ends rounded, not constricted, thick-walled, one-septate, hyaline, $9-14 \times 4.5-6 \mu$.

On decayed stumps and logs, also on *Polyporus squamosus*. Generally distributed.

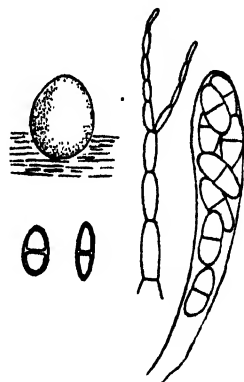


Fig. 9. *Dialonectria galligena*; perithecium, $\times 30$; paraphysis, $\times 225$; ascus, $\times 450$; ascospores, $\times 450$.

Dialonectria sanguinea (Bolt.) Cooke in *Grevillea*, XII (1884), 110; *Sphaeria sanguinea* Bolton, *Fungi Halifax*, III (1789), 121; *Nectria sanguinea* (Bolt.) Fr., *Summa Veg. Scand.* (1849), 388; *Sphaeria Purtoni* Grev. *Scottish Crypt. Flora*, VI (1828), Synopsis, p. 23; *Nectria Purtoni* (Grev.) Currey in *Trans. Linn. Soc.* XXII (1858), 282, fig. 182; *Sphaeria episphaeria* Tode, *Fung. Meckl.* II (1791), 21; *Nectria episphaeria* (Tode) Fr., *Summa Veg. Scand.* (1849), 388.

Perithecia superficial, scattered or gregarious, blood-red, becoming dark red, ovoid or subconoid, apex papillate, often collapsing laterally, smooth, 0.15–0.25 mm. diameter, wall of rather small cells; asci cylindrical, almost sessile, apex truncate, $55-80 \times 4-6 \mu$, spores uniseriate or obliquely uniseriate; ascospores oval or oblong oval, sometimes subfusoid, one-septate, not or slightly constricted, hyaline, becoming pale yellow and minutely granular, $9-14 \times 4-6 \mu$.

On rotten wood and Sphaeriaceous fungi. Generally distributed.

Dialonectria Brassicae (Ellis & Sacc.) Cooke in *Grevillea*, XII (1884), 110; *Nectria Brassicae* Ellis & Sacc. in *Michelia*, II (1881), 374.

Perithecia superficial, scattered or clustered, minute, 0.15–0.18 mm. diameter, blood-red, globoso-conoid or globose with a papillate apex, glabrous, apex hyaline by transmitted light; asci clavate, almost sessile, apex truncate and impressed, $63-75 \times 9 \mu$, spores uniseriate, or biseriate above; no paraphyses; ascospores narrow-oval, often subfusoid, sometimes cymbiform, ends rounded, one-septate, sometimes constricted, hyaline, becoming brownish, $11-18 \times 4-6 \mu$, rarely 9μ long. [Weese's measurement is $8-13 \times 3-4 \mu$.]

On decaying stalks of *Brassica*. North Wootton, October 1935.

Dialonectria Desmazierii (de Not.) Petch, in *Naturalist* (1937), 281; *Nectria Desmazierii* de Not., *Sfer. Ital. Cent.* I, no. 4 (1863); *Sphaeria*

sanguinea var. *cicatricum* Berk. in *Mag. Zool. Bot.* 1 (1837), 48; *Sphaeria coccinea* var. *cicatricum* Desm. in *Ann. Sci. Nat.* ser. 3, x (1848), 351; *Nectria cicatricum* Tul., *Sel. Fung. Carp.* III (1865), 77.

Perithecia scattered, superficial, or clustered on a feebly developed hyphal layer which turns black with age, blood-red, minute, conoid or urceolate, 0.18 mm. high, 0.15 mm. diameter, smooth, apex rounded, often collapsing laterally; asci cylindrical, almost sessile, $72-85 \times 8 \mu$; ascospores ellipsoid or oblong, ends rounded, one-septate, upper cell sometimes the broader, hyaline, becoming pale yellow, minutely warted, $9-15 \times 4.5-6 \mu$.

On twigs of box, commonly on the leaf scars. Apethorpe (Berkeley, *British Fungi*, no. 83) Hb. B.M. and Hb. Kew.; Wiltshire (Broome) Hb. B.M.; Overstrand Woods, October 1934; Buckden, September 1936.

Dialonectria galligena (Bres.) Petch, in *Cat. Yorks. Fungi*, (1937) 32; *Nectria galligena* Bres. in Strasser, *Pilzfl. Sonntagbl.* IV (1901), 413.

Perithecia superficial, scattered or densely crowded, sometimes caespitose, orange or pale red (rarely yellow), becoming dark red and finally blackening, globose to ovate, 200–300 μ diameter, smooth, ostiolum papillate, not collapsing; asci clavate, sessile, apex rounded, $72-92 \times 9-15 \mu$; paraphyses stout, branched, somewhat persistent; ascospores fusoid or oval, ends rounded, one-septate, hyaline, becoming pale yellow, $12-27 \times 4-9 \mu$.

Conidial stage, *Fusarium Wilkommii* Lind., pustules erumpent, white, farinose, linear or circular; conidia cylindric or linear-clavate, straight or slightly curved, one- to five-septate, $36-60 \times 4 \mu$.

Distinguished from scattered forms of *Nectria coccinea* by the rounded apex of the ascus and the stout branched paraphyses.

The cause of "canker" on apple and pear trees. Generally distributed. Also on ash, Charminster, Dorset, December 1894, Hb. B.M.; Chilterns, October 1933 (Ehrlich), Hb. Kew. Originally described from specimens on galls on *Salix purpurea*, and recorded in Europe on beech, hazel, and bird-cherry.

Dialonectria graminicola (B. & Br.) Cooke in *Grevillea*, XII (1884), 110; *Nectria graminicola* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 3, III (1859), 376.

Perithecia superficial, scattered, each on a small, pale brown, basal membrane, globose or ovate, about 0.25 mm. diameter, collapsing and becoming cup-shaped, red, brown when old, smooth, ostiolum papillate, small; asci sessile, fusiform, apex truncate, $50-62 \times 8.5-10 \mu$; ascospores fusoid, ends rounded, one-septate, hyaline, $16-20 \times 3.5-4.5 \mu$.

On dead leaves of *Aira caespitosa*. Batheaston, January 1859 (Broome), Hb. B.M. and Hb. Kew.

Dialonectria arenula (B. & Br.) Cooke in *Grevillea*, XII (1884), 110; *Sphaeria arenula* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, IX (1852), 320; *Nectria arenula* (B. & Br.) Berk., *Outlines* (1860), 394.

Perithecia superficial, usually scattered, ochre-yellow, subglobose or ovate, flattened above, not collapsing, up to 0.3 mm. high, 0.27 mm. diameter, usually with a short, stem-like base, sometimes slightly rough, with a minute, conical ostium; asci clavate, $70 \times 10 \mu$; ascospores fusoid, one-septate, rather strongly attenuated downwards, often inequilateral, straight or curved, or oval or subcylindric, $14-20 \times 4-6 \mu$, ends rounded, usually constricted.

On dead leaves of *Aira caespitosa*. Batheaston, February 1851 (Broome), Hb. B.M. and Hb. Kew. The specimen recorded by Bucknall, on dead leaves of *Iris pseudacorus*, near Aust, June 1881, is *Nectriella dacrymycella* (specimen in Hb. B.M.).

Dialonectria Wegeliana (Rehm) Petch, comb.nov.; *Nectria episphearia* f. *Wegeliana* Rehm in *Hedwigia* (1891), 260; *Nectria Wegeliana* (Rehm) v. Höhn., in Strasser, *Verhandl. zool.-bot. Gesellsch., Wien*, LV (1905), 604.

Perithecia superficial, scattered or clustered, ovoid, apex conico-discoid, 0.18 mm. diameter, 0.24 mm. high, sometimes collapsing, blood-red, blackening, minutely rugose, smooth round the ostium; outer layer of the perithecial wall hyaline, wholly or in part; asci cylindrical or clavate, apex rounded or truncate, sessile, $90-100 \times 9-10 \mu$, spores obliquely uniseriate; paraphyses slender, diffuent; ascospores broadly ellipsoid, ends rounded, one-septate, hyaline becoming brownish and warted, $10-18 \times 6-9 \mu$, with some globose, continuous, 9μ diameter.

On *Diatrypella* on birch, Wraxall, February 1845 (Broome), Hb. Kew.; on *Diatrypella*, Lytchett, October 1922 (Hawley), Hb. B.M.; on *Diatrypella quercina*, Carlisle, Hb. Kew.; ditto, Logan Woods, near Stranraer, Wigtownshire, August 1936 (C. G. C. Chesters).

Dialonectria Veuillotiana (Sacc. & Roum.) Cooke in *Grevillea*, XII (1884), 110; *Nectria Veuillotiana* Sacc. & Roum. in '*Michelia*', II (1881), 325.

Perithecia superficial and densely crowded, urceolate, i.e. globose or ovoid, 0.4-0.5 mm. diameter, 0.5 mm. high, with a flat projecting apical disc, rough with conspicuous warts, yellowish red, becoming scarlet to red-brown, ostium minute, scarcely or not projecting; asci cylindrical or clavate, $80-100 \times 9-12 \mu$, apex rounded or subtruncate, spores uniseriate or obliquely uniseriate; paraphyses delicate, linear, diffuent; spores ellipsoid or oval, often inequilateral, hyaline, one-septate, minutely warted, $14-20 \times 5-7 \mu$. Stout, septate, red hyphae, $14-16 \mu$ diameter, spread over the host from the base of the perithecium.

Ralfs 557 in Hb. B.M., without locality; on beech, Knole Park, October 1933 (Ehrlich), Hb. Kew.

LASIONECTRIA Cooke in *Grevillea*, XII (1884), 112

Perithecia and ascospores as in *Dialonectria*, but perithecia hairy or setose.

Lasionectria flavida (Corda) Cooke in *Grevillea*, XII (1884), 110; *Sphaeria flavida* Corda, *Icon. Fung.* IV (1840), 40, fig. 117; *Nectria flavida* (Corda) Fr., *Summa Veg. Scand.* (1849), 388; *Calonectria flavida* (Corda) Sacc. in *Michelia*, I (1878), 313.



Fig. 10. *Lasionectria lecanodes*; perithecium, $\times 50$; ascospores, $\times 800$.

Perithecia superficial, scattered or gregarious, small, 60–120 μ diameter, globose with a minute, conical ostiolum, at first golden yellow, densely clothed with short, yellow tomentum except at the ostiolum, and with longer yellow hyphae extending in strands from the lower half of the perithecium over the substratum and coalescing with those from adjacent perithecia, the perithecial wall gradually turning red from the apex downwards, old and weathered perithecia being red and more or less naked; asci clavate, sometimes cylindrical, apex subtruncate, 40–60 \times 6–8 μ , spores obliquely uniseriate or biseriate; paraphyses diffluent; ascospores oblong or subfusoid, inequilateral or subcymbiform, sometimes slightly curved, distinctly one-septate, sometimes slightly constricted, 9–15 \times 2.5–3 μ .

On dead wood, Leigh Wood, Bristol, 15 December 1847 (Broome), Hb. Kew. and Hb. B.M.; on bramble, Batheaston, 7 March 1869 (Broome), Hb. Kew. and Hb. B.M.; on sawdust, Langridge, April 1874, Hb. Kew. and Hb. B.M.; on dead wood, Scarborough (Massee), Hb. B.M.; ditto, Missenden, 10 January 1925 (E. J. H. Corner); on decayed wood of beech, Nun Appleton, Yorks, 14 April 1935 (W. G. Bramley).

I have accepted Berkeley and Broome's interpretation of Corda's species, as it appears to fit Corda's description and figure better than any other which has been proposed.

Lasionectria aureola (Winter) Sacc., *Syll. Fung.* IX (1891), 970; *Nectria aureola* Winter in *Hedwigia*, XXIV (1885), 261.

Perithecia gregarious, superficial, minute, globoso-conoid, pale yellow, subtranslucent, covered above with short, rigid, simple, white, spreading hairs; asci oblongo-fusiform, shortly stalked, 47–53 \times 7 μ ; ascospores fusiform, slightly attenuated and narrowly rounded at the ends, hyaline, one-septate, 14 \times 2.5 μ (Winter).

On the mycelium of *Meliola Niessleana* on *Vaccinium Vitis-Idaea*, Killin, Perthshire, July 1907 (D. A. Boyd), Hb. B.M.

Lasionectria lecanodes (Ces.) Petch, comb. nov.; *Nectria lecanodes* Ces. in Rabh. *Herb. Myc.* ed. II, no. 525.

Perithecia superficial, scattered or gregarious, depressed globose, almost discoid when dry, 0.24 mm. diameter, umbilicate, pinkish, clothed with minute, white, erect or decumbent hairs which are up to 40 μ long, 2–3 μ diameter, simple or branched, irregular, con-

torted above and sometimes toothed, with a rounded or globose apex; asci cylindrico-clavate, apex truncate, $60-70 \times 5-6 \mu$; ascospores oval or oblong-oval, ends rounded, hyaline, minutely warted, one-septate, $9-12 \times 3-4 \mu$.

On the thallus of *Peltigera canina*, near King's Lynn, October 1876, 1877, Plowright, *Sphaer. Brit.* III, no. 12, and Cooke, *Fung. Brit. Exsicc.* II, no. 564; North Wootton, 1884 (Plowright) Hb. Kew. On *Lobaria pulmonaria*, Inverary, September 1907 (D. A. Boyd), Hb. B.M.

Lasionectria Leptosphaeriae (Niessl) Petch, comb. nov.; *Nectria Leptosphaeriae* Niessl, in Krieger, *Fung. Sax.* no. 165 (1886).

Perithecia superficial, scattered or in small groups, at first flesh-coloured to brick-red, then dark red, pruinose, globose or broadly conoid, $0.2-0.3$ mm. diameter, with a broad, blackish apical disc bearing a minute, conical ostium, collapsing, furnished above with short, scattered septate hairs with an ellipsoid terminal cell, and below with hyaline hyphae which spread from the perithecium over the substratum; asci narrow-clavate or cylindrical, almost sessile, apex truncate, $90-125 \times 9-13 \mu$, spores obliquely uniseriate; paraphyses narrow strap-shaped, branched, $2-4 \mu$ broad, diffuent; ascospores hyaline or pale yellow, narrow-oval or fusoid, sometimes subcymbiform, one-septate, ends rounded, $15-22 \times 5-6.5 \mu$.

On the perithecia and mycelium of *Leptosphaeria*. On nettle, Batheaston, November 1861 (Broome); on bramble, Charny Down, October 1864 (Broome); Little Eaton, July 1910 (Gibbs); all in Hb. B.M., det. Ehrlich. Staunton, October 1862 (Berkeley), Hb. Kew., det. Ehrlich.

NEOHENNINGSIA Koorders, *Bot. Untersuch.* (1907), 164

Perithecia and ascospores as in *Dialonectria*, but perithecia furnished with a ring of more or less horizontal, long, triangular processes round the apex.

Neohenningsia suffulta (B. & C.) v. Höhnelt, *Fragm. z. Mykol.* no. 755 (1912); *Nectria suffulta* B. & C. in *Journ. Linn. Soc.* X (1868), 378; *Nectria ornata* Masee & Salmon in *Ann. Bot.* XVI (1902), 75, figs. 29-32.

Perithecia superficial, gregarious, orange-yellow, globose, up to 0.5 mm. diameter, flattened above and furnished at the margin of the flattened area and near it with flat, silvery, spreading scales or tufts of hyphae, $50-110 \mu$ long, $15-25 \mu$ broad; asci cylindrical or cylindrico-clavate, $65-75 \times 9-11 \mu$; ascospores ellipsoid, ends obtuse, one-septate, hyaline, $12-14 \times 4-5 \mu$.

On horse dung, Kew. (Masee and Salmon), Hb. Kew.

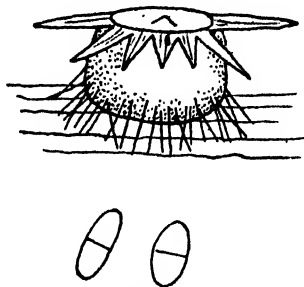


Fig. 11. *Neohenningsia suffulta*; perithecium, $\times 40$; ascospores, $\times 500$.

NECTRIELLA Nitschke, in Fuckel, *Symb. Mycol.* (1869), 175

Perithecia separate, immersed, then erumpent, bright coloured; ascospores as in *Dialonectria*.

Nectriella dacrymycella Rehm, *Ascomyceten*, no. 232 b; Thuemen, *Mycoth.* no. 1064.

Perithecia scattered, subepidermal, erumpent, yellow, brownish yellow or reddish, globose, 300–500 μ diameter, with a minute, papillate ostiolum, collapsing and becoming convex; asci clavate, usually with a long pedicel, 65–95 \times 9–14 μ , spores obliquely uniseriate or biseriate; ascospores hyaline, fusoid, inequilateral, sometimes curved, one-septate, not constricted, 15–22 \times 4–5 μ .

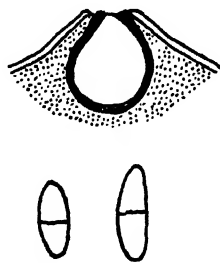


Fig. 12. *Nectriella Robergei*; perithecium in median section, \times 50; ascospores, \times 600.

On *Iris*, near Aust, June 1881 (Bucknall), Hb. B.M.

Phillips and Plowright (*New and Rare British Fungi*, no. 291) recorded *Nectria dacrymycella* (Nyl.) Karst., collected by Bucknall on *Angelica sylvestris*, Blaize Castle Wood, May 1882. The specimen is not in Hb. Kew. or Hb. B.M., and it is not possible to say what it was, as the description given by Phillips and Plowright was taken, except for one line, from Karsten. Several species have passed under that name, and Weese regards Karsten's fungus as *Nectriella luteola* (Rob.) Weese. On the other hand, Bucknall's specimen on *Iris*, referred by him to *Nectria arenula* B. & Br., is not the latter species, but matches *Nectriella dacrymycella* Rehm in Rehm, *Ascomyceten*, no. 232 b.

Nectriella Bloxami (B. & Br.) Fuckel, *Symb. Mycol. Nachtr.* III (1875), 21; *Nectria Bloxami* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, XIII (1854), 467; *Calonectria Bloxami* (B. & Br.) Sacc., *Fung. Veneti*, ser. IV, 23.

Perithecia scattered, subepidermal, erumpent, depressed globose or ovoid, about 0.25 mm. diameter, collapsing, dark red, smooth; asci not seen; ascospores cymbiform, ends broadly rounded, evidently one-septate, 11–16 \times 4–4.5 μ .

On dead stems of Jerusalem artichoke, Twycross (Bloxam), Hb. Kew. and Hb. B.M. The type specimens are in poor condition.

Nectriella funicola (B. & Br.) Petch in *Naturalist* (1937) 281; *Sphaeria funicola* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, VII (1851), 188; *Nectria funicola* (B. & Br.) Berk., *Outlines* (1860), 393; *Calonectria funicola* (B. & Br.) Sacc. in *Michelia*, I (1878), 312; *Nectriella charticola* Fuckel, *Symb. Mycol.* (1869), 176; *Nectria fibricola* Plowright, apud Sacc. in *Michelia*, II (1880), 152; *Lasionectria funicola* (B. & Br.) Cooke in *Grevillea*, XII (1884), 112.

Perithecia immersed, raising and separating the upper layers of the substratum and becoming partly exposed, scattered or in small groups, dull brown or orange, globose up to 0.36 mm. diameter, convex above, or flask-shaped, or globose with a tubular neck up to

0.3 mm. long, 0.1 mm. diameter, naked, or with the exposed part sparsely covered with short, rigid, spreading, white hairs; hairs up to 50μ long, $4-5\mu$ diameter below, hyaline, tapering upwards, thick-walled below, apex obtuse or acute and sometimes shortly encrusted; asci clavate, apex truncate, $90-120 \times 9-20\mu$; ascospores narrow-oval, often inequilateral, or subcymbiform, one-septate, the lower cell sometimes attenuated, ends rounded, hyaline, $14-22 \times 4-6\mu$.

On decayed rope, King's Cliffe, October 1841 (Berkeley), Hb. Kew.; on rotting string, King's Lynn, 1881 (Plowright), Hb. B.M.; on old cardboard, Melton Wood, near Cadeby, September 1901 (Masse and Crossland), Hb. Kew.

Nectriella Robergei (Mont. & Desm.) Weese in *Ann. Mycol.* xii (1914), 138; *Nectria Robergei* Mont. & Desm., *Pl. Crypt. France* (1856), fasc. viii, no. 374; *Nectria lichenicola* (Ces.) Wint. in *Flora* (1872), 523; *Nectria Peltigerae* Phillips & Plowright in *Grevillea*, iv (1876), 123 (223 in error).

Perithecia immersed, scattered, epiphyllous, raising the cortical layer and splitting it stellately, sordid flesh-coloured, pale brownish red, or orange-red, vertically oval, up to 240μ diameter, 270μ high, apex rounded; asci clavate, apex truncate, $66-85 \times 8-13\mu$; ascospores broadly oval, one-septate, wall and septum thin, $10-15 \times 4-7\mu$, containing numerous small oil globules. [According to Weese (*loc. cit.*), the asci in the type of *Nectriella Robergei* are $45-68 \times 9-15\mu$.]

Conidial stage, *Illosporium carneum* Fr.

In the thallus of *Peltigera canina*. North Wootton, 13 November 1873 (Plowright), Hb. B.M. and Hb. Kew.; Castle Rising, 1 September 1875 (Plowright), Hb. B.M.; near King's Lynn, November 1875, Plowright, *Sphaer. Brit.* iii, 13, and Vize, *Microf. Brit.* no. 388. Also recorded on the same host from Forden by Vize.

HYPHONECTRIA Sacc., *Syll. Fung.* ii (1883), 501 (as subgenus) (*Nectriopsis* Maire in *Ann. Mycol.* ix (1911), 323)

Perithecia superficial, separate, on a common byssoid subiculum, bright-coloured, soft; asci eight-spored; ascospores hyaline, one-septate, not apiculate.

Hyphonectria muscivora (B. & Br.) Petch, comb. nov.; *Nectria muscivora* (B. & Br.) Berk., *Outlines* (1860), 394; *Sphaeria muscivora* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, vii (1851), 188; *Calonectria muscivora* (B. & Br.) Sacc. in *Michelia*, i (1878), 315; *Sphaeria bryophila* Rob., in Desm. in *Ann. Sci. Nat.* ser. 3, xvi (1851), 306; *Nectria bryophila* (Rob. & Desm.) Sacc. in *Michelia*, i (1878), 296.

Perithecia seated on, or partly embedded in, a white floccose subiculum, scattered or clustered, oval or pyriform,

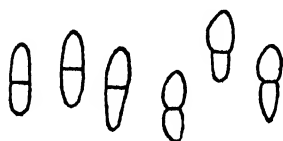


Fig. 13. *Hyphonectria aureonitens*; perithecia, $\times 30$; ascospores, $\times 800$.

with a prominent papilla, up to 0.35 mm. high, 0.28 mm. diameter, bright orange, becoming amber when dry; asci clavate, sessile, $80-100 \times 12-14 \mu$; ascospores narrow-oval to subfusoid, sometimes subcymbiform, ends rounded or subacute, one-septate, $15-24 \times 6-8 \mu$.

On mosses on the mud tops of walls in winter, King's Cliffe (Berkeley). Also recorded from Dolgelly (B.M.S.). The record from Hebden Bridge, Yorks, is incorrect.

Hyphonectria violacea (Schmidt) Petch in *Journ. Bot.* LXXV (1937), 220; *Sphaeria violacea* Schmidt, in Fries, *Syst. Myc.* II (1822), 441; *Hypomyces violaceus* (Schm.) Tul. in *Ann. Sci. Nat.* ser. 4, XIII (1860), 14; *Peckiiella violacea* (Tul.) Sacc., *Syll. Fung.* IX (1891), 945; *Nectriopsis violaceus* (Schm.) Maire in *Ann. Mycol.* IX (1911), 323; *Byssonectria violacea* (Schm.) Seaver in *Mycologia*, II (1910), 65; *Hypomyces candicans* Plowr. in *Grevillea*, XI (1882), 50; *Nectriopsis candicans* (Plowr.) Maire in *Ann. Mycol.* IX (1911), 323.

Subiculum white, becoming violet; perithecia partly immersed, or wholly immersed except the ostium, globose, with a small conical ostium, up to 0.2 mm. diameter, wall hyaline to purple-violet; asci cylindrical, $50-60 \times 3-4 \mu$, spores uniseriate or obliquely uniseriate; ascospores oval or oblong-oval, ends rounded, hyaline, smooth, distinctly one-septate when mature, $6-9 \times 2-3 \mu$.

Conidia oval, or linear-oblong, ends obtuse, continuous, sometimes one-septate, $10-22 \times 5-7 \mu$ (Tulasne).

Chlamydospores broadly ovate, or oval, $21-30 \times 16-25 \mu$, or globose, $22-24 \mu$ diameter, yellow-brown, thick walled, smooth.

On *Fuligo septica*. Cawdor Castle, September 1879 (Plowright); Leziate, August 1880 (Plowright); Bathford Down, October 1880 (Plowright), Hb. B.M.

Hyphonectria Berkeleyana (Plowr. & Cooke) Petch in *Journ. Bot.* LXXV (1937), 220; *Hypomyces Berkeleyanus* Plowr. & Cooke in *Grevillea*, XI (1882), 48; *Nectriopsis Berkeleyanus* (Plowr. & Cooke) Maire in *Ann. Mycol.* IX (1911), 323.

Subiculum rose-coloured or pallid; perithecia scattered or gregarious, globose, 0.24 mm. diameter, with a papillate ostium, scarlet, smooth; asci cylindrical, apex truncate, $66-80 \times 5 \mu$; ascospores uniseriate or obliquely uniseriate, oblong-oval, ends rounded, one-septate, smooth, hyaline, becoming pale yellowish and faintly warted, $7-11 \times 4 \mu$.

On *Stereum hirsutum*, Downton, Herefordshire, October 1878 (Plowright). On *Corticium* sp., Sandringham, November 1878 (Plowright). On *Polyporus* sp., Hb. B.M., sub *Nectria episphaeria*, ex Hb. Bloxam.

Hyphonectria aureo-nitens (Tul.) Petch in *Journ. Bot.* LXXV (1937), 220; *Hypomyces aureo-nitens* Tul., *Sel. Fung. Carp.* III (1865), 64; *Nectriopsis aureo-nitens* (Tul.) Maire in *Ann. Mycol.* IX (1911), 323.

Subiculum golden yellow or white, scanty; perithecia minute,

ovoid, 0.15–0.18 mm. high, 0.12–0.15 mm. diameter, golden yellow or pallid, sparsely clothed with yellow or white hyphae, or almost naked, ostiolum punctate, darker; asci cylindrical or narrow clavate, $90 \times 4\text{--}7\ \mu$, spores uniseriate or obliquely uniseriate; ascospores oval, narrow-oval, or fusoid, ends rounded or one end subacute, hyaline, minutely warted, $9\text{--}13 \times 3\text{--}4\ \mu$, ultimately becoming constricted at the septum, with the upper cell usually the broader.

Conidial stage, *GlIOClaDIum penicillioides* Cda, *Icon. Fung.* iv, 31; *Verticillium Aspergillus* B. & Br., *Notices of British Fungi*, no. 1384. Conidiophores clustered in small scattered white tufts; stalk stout, $5\text{--}6\ \mu$ diameter, rigid, minutely verrucose, about $200\ \mu$ high, branched in the upper half or third, thickened at the point of origin of the branches; primary branches opposite, sometimes solitary, stout, cylindrical, usually about $20\ \mu$ long, but sometimes longer and septate, curving upwards parallel to the main stem; secondary branches in whorls of two to four, cylindrical, about $10\ \mu$ long, each bearing an apical cluster of three to five, narrow flask-shaped or conoid phialides of about the same length, all erect, parallel and crowded; conidia oval or subcylindrical, ends obtuse, hyaline, $2\text{--}5 \times 1.5\text{--}2\ \mu$, rarely $7 \times 2\ \mu$, collected in an apical globule supported by all the phialides.

On *Stereum hirsutum*, Pwll-y-crochan Wood, North Wales, 11 October 1880 (Plowright), Hb. Kew. and Hb. B.M.; on *Stereum*, Hubberholme, Yorks, 9 September 1936; on *Stereum*, Keld, Yorks, October 1936 (W. G. Bramley).

HypHonectria Solani (Reinke & Berth.) Petch in *Journ. Bot.* LXXV (1937), 220; *Hypomyces Solani* Reinke & Berth., *Zersetz. d. Kartoffel* (1879), 27.

Subiculum loose, byssoid; perithecia pale purple-red, orange-yellow at the apex, obpyriform; ascospores oblong-oval, ends subacute, one-septate, constricted at the septum, verrucose, $13\text{--}16 \times 7\text{--}8\ \mu$.

On rotting potatoes. Ireland (Pethybridge in *Trans. Brit. Mycol. Soc.* vi (1918) 11).

HYPOMYCES Tulasne in *Ann. Sci. Nat.* ser. 4, xiii (1860), 11 (in part)

Perithecia superficial, separate, on a common byssoid subiculum, bright-coloured, soft; asci usually eight-spored; ascospores hyaline, one-septate, apiculate.

Hypomyces ochraceus (Pers.) Tul. *loc. cit. supra*, p. 12; *Sphaeria ochracea* Pers., *Synopsis* (1801), 18; *Hypocrea apiculata* Peck in *Rep. N.Y. State Mus.* xxix (1878), 57; *Hypomyces apiculatus* (Peck) Seaver in *Mycologia*, II (1910), 73; *Hypomyces terrestris* Plowr. & Boud. in *Grevillea*, viii (1880), 105.

Subiculum on the ground, where the affected agaric has decayed, white, then flesh-coloured or peach-coloured, becoming ochraceous

when old and forming compact patches, 1–2 cm. diameter, resembling a *Corticium*; perithecia at first immersed in the stroma, then nearly half-free, crowded, conoid, or subglobose with a conoid apex, 0.3 mm. diameter, yellow or reddish; asci cylindrical, $150-200 \times 6-7 \mu$; ascospores lanceolate or narrow-oval, often inequilateral, apiculate, verrucose, one-septate, becoming constricted when old, hyaline, $27-40 \times 6-7 \mu$.

Conidial stage, *Verticillium agaricinum* Corda (*V. Lactarii* Peck), on the affected agaric, white, effused, even, becoming floccose; conidiophore verticillioïd, with branches in whorls of two to four, but often solitary; tapering to the apex; conidia apical, solitary, rarely two or three together, oval, obovate, or oblong, with a broad, truncate apiculus, continuous, $11-21 \times 9-12 \mu$, or globose, $10-13 \mu$.

Chlamydospores, *Blastotrichum puccinioides* Preuss, produced on the deeper hyphae of the conidial stage, compound, fusoid or narrow-oval, two- to four-septate, strongly constricted, the central cell the largest, apical cell rounded or conoid, $70-140 \times 24-33 \mu$, or oblong-oval, one-septate, $48 \times 28 \mu$, hyaline, becoming reddish purple, smooth (? finally minutely warted), often budding from the central cell.

On *Russula* and *Lactarius*, the conidial and chlamydospore stages on the agaric, but the perithecial stage on the ground, or on vegetable debris, after the agaric has decayed. North Wootton, 1872–80 (Plowright), Hb. Kew.; Leziate, 1880 (Plowright); Ashwick, Middleton, October 1896 (Plowright), Hb. Kew.; Haslemere, September 1930 (Swanton), Hb. Kew.; Midhurst (Miss Wakefield), Hb. Kew.; Epping Forest, September 1936 (A. A. Pearson). The identity of *H. apiculatus* and *H. terrestris* was determined by Miss Wakefield.

Hypomyces rosellus (A. & S.) Tul., *loc. cit. supra*, p. 12; *Sphaeria rosella* A. & S., *Consp. Fung.* (1805), 38, pl. 7, fig. 3; *Nectria Albertini* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 3, vii (1861), 452.

Subiculum white, then rose-coloured; perithecia gregarious, globoso-conoid, 0.15 mm. diameter, deep rose-red; asci cylindrical, $150 \times 6.5 \mu$; ascospores lanceolate, straight or curved, apiculate, apiculus sometimes curved, hyaline, one-septate, slightly constricted, verrucose, $22-37 \times 5-7 \mu$.

Conidial stage, *Dactylium dendroides* Fr., conidiophore verticillioïd; conidia cylindrico-oblong, obtuse, one- to three-septate, $25-35 \times 10-13 \mu$.

On decaying fungi, most frequently *Stereum hirsutum*. Generally distributed. Plowright, *Sphaer. Brit.* iii, no. 4.

Hypomyces aurantius (Pers.) Tul., *Sel. Fung. Carp.* iii (1865), 48; *Sphaeria aurantia* Pers., *Synopsis* (1801), 68; *Nectria aurantia* Fr., *Summa Veg. Scand.* (1849), 388; *Sphaeria aurea* Grev., *Scottish Crypt. Flora*, pl. 47 (in part).

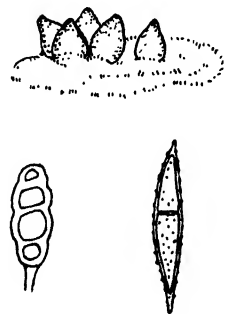


Fig. 14. *Hypomyces aurantius*; perithecia, $\times 15$; chlamydospore, $\times 250$; ascospore, $\times 800$.

Subiculum floccose, ochraceous, often with a white margin; perithecia crowded, globose with a conoid apex, up to 0.3 mm. diameter, 0.36 mm. high, golden yellow or orange, becoming orange-red, then dark red, yellow pruinose; asci cylindrical, $110-140 \times 6 \mu$; ascospores lanceolate, acute, apiculate, usually curved, hyaline, verrucose, one-septate, $18-27 \times 4-6 \mu$.

Conidial stage, *Diplocladium penicillioides* Sacc., conidiophores with a main stem, 4μ diameter, bearing flask-shaped or conoid phialides, about 30μ long, in whorls of three at the apex, opposite or solitary elsewhere, with terminal, solitary conidia; conidia oval or oblong-oval, one end apiculate, one-septate, $9-16 \times 6-9 \mu$, hyaline.

Chlamydospores compound, oblong-oval, one- to three-septate, constricted at the septa, pale brown or reddish brown, thick-walled, smooth, $18-48 \times 12-18 \mu$.

On *Polypori* and the tougher *Agaricini*, *Polyporus squamosus*, *P. picipes*, *P. adustus*, *Polystrictus versicolor*, *Panus torulosus*, etc. Generally distributed. Berkeley and Broome noted that a very pale, honey-coloured variety, springing from a snow-white subiculum, sometimes accompanied the darker form. There is a specimen of this in Hb. Kew., collected by J. Harvey Bloom at Mickleham, November 1927. In the United States, this form has been named *Hypomyces polyporinus* Peck.

***Hypomyces Broomeanus* Tul., Sel. Fung. Carp. III (1865), 108.**

Subiculum white, or pale brown, compact; perithecia ovate, 0.25-0.3 mm. diameter, apex subacute or sometimes produced, hyaline becoming pale brown, densely clothed with white or pale brown, short tomentum, except at the apex, superficial, or immersed through overgrowth of the tomentum; asci cylindrical, $130-140 \times 4 \mu$; ascospores fusoid, one-septate, coarsely warted, usually shortly apiculate at each end, hyaline, $13-16 \times 3.5-4 \mu$.

Conidial stage, *Gliocladium strictum*, n.sp., conidiophores crowded, white in mass, hyaline, about 100μ high; stem stout, 4μ diameter below, with few branches, the lower branches usually solitary and distant, the upper branches (phialides) opposite or in whorls of three, about 30μ long, tapering, all branches parallel to the main stem; conidia hyaline, oblong-oval or narrow-oval, slightly inequilateral, ends obtuse, $5-11 \times 2-2.5 \mu$, the longer ultimately becoming one-septate or pseudo-septate, united by mucus in heads about 14μ diameter, and ultimately in large masses.

On *Fomes annosus*. Ashton Court, January 1845 (Broome), Hb. B.M.; Staunton, Notts, September 1851 (Berkeley), Hb. Kew. and Hb. B.M.; Lucknam, December 1864 (Broome), Hb. B.M.; Bathford, November, December 1864 (Broome), Hb. B.M. and Hb. Kew., Rabh. *Fung. Europ.* no. 751; Batheaston, 5 November 1877 (Broome), Hb. B.M.; King's Lynn, November 1875, 1876 (Plowright), Hb. Kew. and Hb. B.M., Plowr. *Sphaer. Brit.* III, 5; North Wootton, March 1936; Hubberholme, September 1936.

***Hypomyces asterophorus* Tul., Sel. Fung. Carp. III (1865), 55.**

Perithecia crowded among *Nyctalis* chlamydospores, globoso-ovoid, conical above, with a ciliate ostium, 150μ high, $70-90 \mu$ diameter,

hyaline, becoming pale yellow-brown; wall composed of large polygonal cells; asci broadly ovate, four- to six-spored, $40-50 \times 18-20 \mu$; ascospores lanceolate, curved, apiculate, one-septate, pale yellow-brown, $25-35 \times 6 \mu$.

Conidial stage, *Polyscytalum fungorum* Sacc., conidiophore branched; conidia cylindrical, hyaline, catenulate, $10-15 \times 3.5 \mu$.

On *Nyctalis*. Hockering Wood, Norfolk, September 1880 (Plowright).

APIOCREA Sydow in *Ann. Mycol.* xviii (1920), 186

Perithecia as in *Hypomyces*; ascospores very unequally one-septate.

Apiocrea chrysosperma (Tul.) Syd. in *Ann. Mycol.* xviii (1920), 186; *Hypomyces chrysospermus* Tul. in *Ann. Sci. Nat.* ser. 4, xiii (1860), 16.

Subiculum golden yellow (from the chlamydospores); perithecia crowded, conoid, up to 0.36 mm. diameter, 0.3–0.32 mm. high, hyaline, becoming orange-yellow, red-brown when dry, smooth; asci cylindrical, or broadly clavate, $120-200 \times 4-10 \mu$; ascospores lanceolate, apiculate at one or both ends, hyaline, very unequally two-celled, one cell sometimes appearing as a small appendage, $21-30 \times 5-6 \mu$, or almost equally two-celled, narrow oval or fusoid, $9-16 \times 3-4 \mu$, rough.

Conidial stage white, effused; conidia pyriform, or oblong-oval, slightly contracted in the middle, continuous, becoming one- or two-septate, $10 \times 5 \mu$ to $30 \times 12 \mu$; conidiophores irregularly verticilloid, often clustered.

Chlamydospores, *Sepedonium chrysospermum* Link, spherical, verrucose, golden yellow, becoming yellow-brown, $10-26 \mu$ diameter.

On *Boleti*, *Paxillus involutus*, *Scleroderma*. Conidial and chlamydospore stages common, perithecia rare. Perithecial stage, Coed Coch (Berkeley), Hb. Kew.; Chapelton Wood near Forres (Stevenson and Plowright), Hb. B.M.; Dartington, Devon (B.M.S.).

Apiocrea Tulasneana (Plowr.) Petch in *Journ. Bot.* Lxxv (1937), 220; *Hypomyces Tulasneanus* Plowright in *Grevillea*, xi (1882), 46, pl. 152, fig. 1; *Hypomyces luteovirens* Tul., *Sel. Fung. Carp.* iii (1865), 57, pl. 8, figs. 15, 16; *Sphaeria luteovirens* Fr., *Syst. Mycol.* ii (1822), 339 (in part); *Peckiella Tulasneana* (Plowr.) Sacc., *Syll. Fung.* ix (1891), 944.

Subiculum sordid yellowish green; perithecia conoid, or subglobose with a cylindrical apex, 0.36–0.5 mm. high, 0.3 mm. diameter, wall yellow, sometimes green at the apex; asci cylindrical, $120-150 \times 10 \mu$;

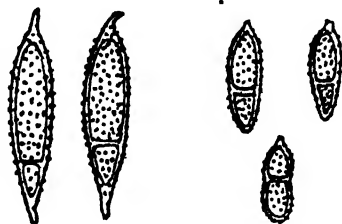


Fig. 15. *Apiocrea chrysosperma*; ascospores of the large-spored and small-spored forms, $\times 1000$.

fusoid, ends rounded, one-septate, generally constricted at the septum and unequally septate, thin-walled, smooth, pale brown, $12-15 \times 4.5-6 \mu$.

Parasitic on *Helminthosporium appendiculatum* (?) and on other species of *Helminthosporium*. Batheaston, January 1859 (Broome), Rabenhorst, *Fung. Europ. Exsicc.* no. 47; Forden, February 1878 (Vize), Plowright, *Sphaer. Brit.* III, no. 10.

CALONECTRIA de Not., *Comm. Critt.* II (1867), 477

Perithecia superficial, separate, bright-coloured (not blue); ascospores oblong, oval, or fusoid, hyaline, two- or few-septate.

Calonectria erubescens (Rob.) Sacc. in *Michelia*, I (1878), 309; *Sphaeria erubescens* Rob. in Desm. *Ann. Sci. Nat.* ser. 3, VI (1846), 72.

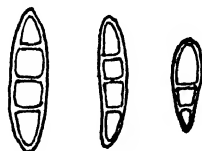


Fig. 18. *Calonectria tessellata*; ascospores $\times 500$.

Perithecia scattered or in small groups, superficial, sometimes with a small rosette of white hyphae at the base, rose-coloured, then brick-red, finally brownish red, globose, about 0.3 mm. diameter, with a minute, papillate ostiolum, almost smooth, collapsing; asci fusoid, $42-65 \times 8-12 \mu$; ascospores hyaline, fusoid or cymbiform, straight or curved, three-septate, $16-22 \times 3-4 \mu$.

On holly leaves. Clifton Downs, October 1879 (Bucknall), Hb. Kew.; Saltash (Rev. J. H. Adams).

Calonectria ochraceo-pallida (B. & Br.) Sacc., *Fungi Veneti*, ser. iv, 23 (1875); *Sphaeria ochraceo-pallida* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, VII (1851), 187; *Nectria ochraceo-pallida* (B. & Br.) Berk., *Outlines* (1860), 394; *Nectria ochraceo-pallida* var. *corallina* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, XIII (1854), 467; *Calonectria Plowrightiana* Sacc. in *Michelia*, I (1878), 307; *Nectria Plowrightiana* (Sacc.) Phil. & Plowr. in *Grevillea*, VIII (1880), 105.

Perithecia scattered or gregarious, superficial, ochre-yellow, sometimes orange-red or pale red, globose, up to 0.35 mm. diameter, slightly flattened above, with a minute papillate ostiolum, slightly darker round the ostiolum, smooth; asci clavate or fusiform, $80-100 \times 14-22 \mu$, spores obliquely uniseriate, or biseriate, or more or less in a parallel bundle; the smaller spores cymbiform, the longer cylindric or fusoid, straight or slightly curved, obtuse, hyaline, three- to eight-septate, usually $22-50 \times 4-6 \mu$, sometimes up to $90 \times 5 \mu$.

On elm, Rockingham Forest (Berkeley), Hb. B.M.; on elder, Gopsall (Bloxam), Hb. B.M.; on elm, King's Cliffe, January 1853 (Berkeley), Hb. Kew.; on hawthorn, Batheaston, 26 February 1877 (Broome), Hb. B.M.; on elm, Mossburnford (A. Jerdon), Hb. Kew.; on poplar, Penzance, Hb. Kew.; on beech, Carlisle, Hb. Kew.; on dead stems of *Arctium Lappa*, near Shrewsbury, February 1878 (Phillips and Plowright), Plowright, *Sphaer. Brit.* III, no. 15; on *Arctium Lappa*, Mulgrave Woods, September 1912, Hb. B.M.; on elm, King's Lynn, Plowright, *Sphaer. Brit.* III, no. 9; Cooke, *Fung. Brit. Exsicc.* ser. I, no. 665.

***Calonectria tessellata* Petch, n.sp.**

Perithecia scattered or crowded, superficial, dull orange when moist, brownish yellow when dry, conoid or ovoid, up to 0.3 mm. high, 0.25 mm. diameter, sometimes collapsing centrally, minutely rugose, glabrous, or with a few projecting cells or small warts at the apex, ostiolum naked, slightly darker; wall with an outer layer of large cells which give the perithecium a rugose or tessellated appearance under a medium magnification, brownish yellow by transmitted light; asci clavate, eight-spored, sessile, apex truncate, $75-95 \times 12-18 \mu$, spores obliquely uniseriate or biseriate; ascospores hyaline, three-septate, oval or oblong-oval, sometimes inequilateral, straight, ends rounded, $18-26 \times 7-9 \mu$.

On decaying stalks of *Brassica*, North Wootton, November 1935; on dead apple twig, Camberley, November 1920, Hb. B.M.

Differs from *C. ochraceo-pallida* in its shorter, broader ascospores, and in the structure of the perithecium wall.

***Calonectria minutissima* Grove in *Journ. Bot.* LXVIII (1930), 31.**

Perithecia almost globose, $80-100 \mu$ diameter, somewhat pallid, very translucent, almost glabrous, with a small papilla, thin-walled; asci lanceolate, attenuated above and below, usually curved, $40 \times 5 \mu$, spores eight, sometimes fewer, biseriate; spores hyaline, fusoid, bent or curved, either end acute or one end subobtuse, $20-22 \times 1.5 \mu$, continuous, containing seven to nine guttae (*ex* Grove, *loc. cit.*).

On dead culms of *Heleocharis palustris* in marshes near Aldrige, Staffs, September (Grove).

***Calonectria platasca* (Berk.) Sacc. in *Michelia*, 1 (1878), 308; *Sphaeria platasca* Berk., *Eng. Fl.* v (1836), 263; *Nectria platasca* Berk., *Outlines* (1860), 393.**

"Scattered, perithecia orange globose confluent with the sub-obtuse ostiolum, base immersed. On the soft wet decayed stump of a maple which had been broken off. Winter. Rockingham Forest, Norths. Rev. M. J. Berkeley.—Perithecia globose, but tapering above into the ostiolum, which varies somewhat in length, so as to have a slightly ovate appearance, immersed in the soft white wood almost to the base of the ostiolum, of the same colour as *Peziza aurantia*, with now and then a few indistinct filaments. Asci broad above, like those of the following species [*Sphaeria affinis* Grev.]. Sporidia oblong, divided into four articulations, each containing a nucleus" (Berkeley, *loc. cit.*).

Nothing further is known about this species. The type in Hb. Kew., from Morehay Lawn, Northants, does not appear to contain any perithecia now. The description suggests a *Cesatiella*, but Berkeley labelled his herbarium specimens, *Nectria Peziza* var. *platasca*.

GIBBERELLA Sacc. in *Michelia*, I (1877), 43

Perithecia superficial, separate, caespitose on an erumpent, parenchymatous stroma, or scattered, soft, appearing brown, dark blue, or black, but the perithecial wall blue or violet by transmitted light; ascospores hyaline (sometimes pink in mass), ovoid or fusoid, two or few septate (usually three- to five-septate).

Gibberella pulicaris (Fr.) Sacc. in *Michelia*, I, (1877) 43; *Sphaeria pulicaris* Fr. in Kunze & Schm., *Mycol. Hfte.* II (1823), 37.

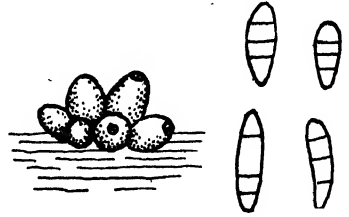


Fig. 19. *Gibberella pulicaris*; perithecia, $\times 25$; ascospores, $\times 500$.

Perithecia caespitose in pulvinate or irregular heaps on a brown stroma, or scattered and solitary, up to 0.3 mm. diameter, globose, conoid, or ovoid, collapsing, at first rufous, then appearing black, but the wall dark violet-blue by transmitted light, rugose, wall leathery, wrinkled and appearing warted when dry, ostiolum minute, papillate; asci subcylindrical or clavate, almost sessile, $75-105 \times 11-15 \mu$, apex at first truncate, becoming rounded, spores obliquely uniseriate or biseriate; ascospores oblong-oval or narrow-oval, sometimes fusoid, inequilateral, straight or curved, ends rounded, three-septate, hyaline, smooth, slightly constricted at the septa, $16-24 \times 6-9 \mu$.

On branches and stems of woody and herbaceous plants. Generally distributed. On elder, broom, and *Brassica* stalks, North Wootton. On *Pteris*, Lyndhurst, August 1885, Hb. Kew.

Gibberella cyanogena (Desm.) Sacc., *Syll. Fung.* II (1883), 555; *Sphaeria cyanogena* Desm. in *Ann. Sci. Nat.* ser. 3, X (1848), 352; *Gibberella Saubinetii* (Mont.) Sacc. in *Michelia*, I (1879), 513; *Gibbera Saubinetii* Mont., *Syll. Crypt.* (1856), 252; *Sphaeria Saubinetii* Dur. & Mont. in *Fl. Alger.* I (1869), 479.

Perithecia scattered, or crowded in flat groups on an inconspicuous brown stroma, up to 0.25 mm. diameter, conoid, or subcylindric, or ovoid and contracted below, collapsing, at first green by transmitted light when fresh, then appearing black, but the wall at first pale blue, then violet-blue, by transmitted light, rugose, covered with large cellular warts, wall thin, ostiolum minute, papillate; asci broadly clavate, with a stout pedicel, $70-100 \times 11-18 \mu$, apex at first truncate, then rounded or acuminate, spores obliquely uniseriate, then biseriate; ascospores narrow-oval, fusoid, or cymbiform, ends rounded, straight or slightly curved, three-septate, rarely four- or six-septate, hyaline, smooth, $20-36 \times 5-7 \mu$, rarely 8μ broad, becoming inflated between the septa on germination.

On decaying herbaceous stems. Common on decaying stalks of *Brassica*. On *Brassica*, Batheaston, October 1855 (Broome), Hb. B.M.; etc. Also on elm, Batheaston, April 1854 (Broome), Hb. B.M.; on broom, Norwich, October 1934; on elder, North Wootton, November 1935.

Gibberella Zeae (Schw.) Petch in *Ann. Mycol.* xxxiv (1936), 260; *Sphaeria Zeae* Schw., *Fung. Car. Super.* no. 234 (1822); Fries, *Syst. Mycol.* II (1823), 527, and III (1829), 232.

Perithecia minute, scattered, or crowded in small, flat groups, usually not united, superficial, without a stroma, subglobose, or ovoid, or conoid, up to 0.25 mm. high, 0.18 mm. diameter, sometimes attached by a broad base, sometimes by a central point, collapsing, appearing black, but the wall violet-grey or smoky blue by transmitted light, bearing, usually on the upper half, rather loose, scattered, small, cellular warts, wall thin, ostium minute, papillate; asci clavate, $55-90 \times 9-13 \mu$, apex truncate, then rounded, spores obliquely uniseriate, then irregularly biseriate; ascospores narrow-oval or cymbiform, straight or slightly curved, three-septate, hyaline, smooth, $18-27 \times 4-5 \mu$.

On stems, etc., of Gramineae, especially cereals. Cockle Park, Northumberland (Dr F. T. Bennett).

Gibberella Buxi (Fuckel) Wint. in *Rabh. Krypt.-Fl.* II (1887), 103; *Gibbera Buxi* Fuckel, *Symb. Mycol. Nachtr.* II (1873), 32; *Lisea Buxi* (Fuckel) Sacc., *Syll. Fung.* II (1883), 518.

Perithecia solitary or in small groups, superficial, conoid, 0.24 mm. high, 0.2 mm. diameter, sometimes collapsing and becoming cup-shaped, smooth, appearing black, but the wall dark violet by transmitted light; asci sessile, broadly clavate, $66 \times 15 \mu$; ascospores oval or oblong-oval, ends rounded, hyaline, one- or three-septate, slightly constricted, $13-20 \times 6-8 \mu$.

On twigs of *Buxus*, with *Dialonectria Desmazierii*, Overstrand Woods, October 1934.

Gibberella acervalis (Moug.) Sacc. in *Michelia*, I (1878), 318; *Sphaeria acervalis* Mougeot, in Fries, *Elench. Fung.* II (1828), 83.

Perithecia caespitose in small groups on an erumpent stroma; stroma plectenchymatous or byssoid, subepidermal, then erumpent; perithecia black, minutely rugose, globose, ovoid or conoid, about 0.25 mm. diameter, collapsing centrally; wall thin, with thicker areas in the upper part, purple-brown or purple-violet by transmitted light; asci cylindrico-clavate, sessile, $80 \times 9-13 \mu$; ascospores narrow-oval or oblong-oval, ends rounded, usually symmetrical, three-septate, $14-22 \times 6-7 \mu$, or ellipsoid or narrow-oval, one-septate, $11-18 \times 5-6 \mu$.

On dead branches of *Salix caprea*, Shelton, Norfolk, February 1873 (Plowright), Hb. B.M. Plowright's specimen is immature, but it appears to be correctly named.

ACTINIOPSIS Starbäck in *Bih. K. Svensk. Vet.-Akad. Handl.* xxv, Afd. III (1899), 54

Perithecia with an apical disc, fringed with spreading triangular fascicles of hyphae, as in *Neohenningsia*; ascospores three- or more septate.

Actiniopsis peristomialis (B. & Br.) Petch, comb. nov.; *Peziza* (*Mollisia*) *peristomialis* B. & Br. in *Ann. Mag. Nat. Hist.* ser. 3, xviii (1866), 126; *Cyathicula peristomialis* (B. & Br.) Sacc., *Syll. Fung.* viii (1889), 308; *Peristomialis Berkeleyi* Boud., *Discom. d'Europe* (1907), 116.

Perithecia at first globose, then subcylindrical, ovoid, or pyriform, apex truncate, 300–400 μ high, 200 μ diameter, pallid, furnished at the margin of the apical disc with long, white, triangular teeth, wall stout; asci subcylindrical or clavate, 80–90 \times 10 μ , spores obliquely uniseriate, then biseriate; paraphyses stout, apex thickened, diffuent; ascospores hyaline, fusiform, inequilateral, subacute, three to five-septate, 20–27 \times 4–5 μ .

On dead bark of holly, Penzance (J. Ralfs), Hb. Kew.

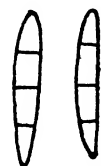


Fig. 20. *Actiniopsis peristomialis*; ascospores, \times 600.

PARANECTRIA Sacc. in *Michelia*, I (1878), 317

Perithecia membranous, subpapillate; ascospores hyaline, three-septate, each end produced into a cilium.

Paranectria affinis (Grev.) Sacc., *loc. cit.*; *Sphaeria affinis* Grev., *Scottish Crypt. Flora*, pl. 186, fig. 1 (1826); *Nectria affinis* (Grev.) Cooke in *Mycologia Scotica* (1879), 362 and *Grevillea*, viii, 9.

Perithecia superficial, scattered or clustered, orange-coloured, globose, 0.25–0.3 mm. diameter, attached by white mycelium from which a few lax hyphae extend over the perithecium, membranous, wall about 20 μ thick, ostiolate, the ostiolum not, or slightly, elevated; asci at first oblongo-clavate, with spores obliquely uniseriate, 40–45 \times 12–14 μ , becoming ovate, 18 μ diameter, with spores in a parallel bundle, eight-spored; ascospores hyaline (or pale yellow?), three-septate, fusoid, sometimes inequilateral, the central part 24–34 \times 6–8 μ , attenuated at each end into a slender seta up to 15 μ long, the upper usually curved, the lower straight.

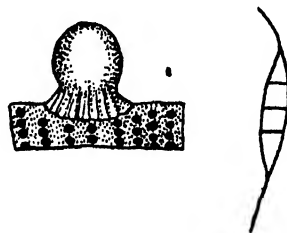


Fig. 21. *Paranectria affinis*; perithecium on the lichen thallus (after Greville), \times 30; ascospore, \times 400.

On *Ephebe lanata* Wainio. Appin (Carmichael), Hb. Kew.

CESATIELLA Sacc., *Michelia*, 1 (1878), 250

Perithecia immersed, separate, soft, pale coloured, ostiolum papillate; ascospores hyaline, two- or more septate.

Cesatiella lancastriensis Grove in *Journ. Bot.* LXVIII (1930), 132.

Perithecia scattered or gregarious, immersed, erumpent, about $500\ \mu$ diameter, colourless, glabrous, soft, narrowly ovoid, apex papillate, obtuse; asci cylindrical or clavato-fusoid, long-stalked, $100\text{--}160(+)\times 12\ \mu$, spores biseriate; ascospores hyaline, fusoid, acute, five-, perhaps seven-septate, $35\text{--}44\times 4\text{--}5\ \mu$; paraphyses very numerous, filiform (Grove, *loc. cit.*).

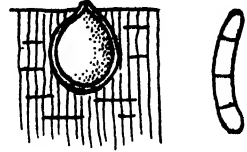


Fig. 22. *Cesatiella*; perithecium in section; ascospore (diagrammatic, after Saccardo).

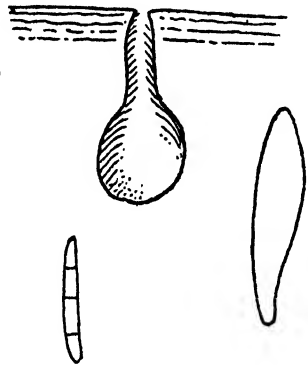
In wet, rotting wood covered with algae, Ainsdale, Lancs, September–December 1928 (Grove).

ORCADIA Sutherland in *Trans. Brit. Mycol. Soc.* v (1914), 151

Perithecia immersed, separate, soft, pallid, with a long beak; ascospores hyaline or yellowish, cylindrical, three-septate.

Orcadia Ascophylli Sutherland, *loc. cit. supra*, pl. 3, fig. 3.

Perithecia entirely immersed, spherical, $160\text{--}200\ \mu$ diameter, with a long, wide, cylindrical beak, $80\text{--}125\ \mu$ long, $35\ \mu$ wide; asci clavate, $65\text{--}75\times 8\ \mu$; paraphyses few, linear, apex slightly inflated; ascospores pale yellow, cylindrical, slightly curved, three-septate (dimensions not given).



In the thallus of *Ascophyllum nodosum*, Orkney.

Fig. 23. *Orcadia Ascophylli*; perithecium (in section), ascus, and ascospore (after Sutherland.)

Orcadia pelvetiana Sutherland in *New Phytol.* xiv (1915), 183, fig. 1.

Perithecia entirely immersed, globose, $110\text{--}140\ \mu$ diameter, with a cylindrical or tapering beak, $160\text{--}180\ \mu$ long, $20\text{--}30\ \mu$ wide; asci clavate, or fusiform and curved, $50\text{--}65\times 11\text{--}13\ \mu$; paraphyses diffuent; ascospores fusiform, curved, about $40\times 4\text{--}5\ \mu$, three-septate, yellowish.

In the thallus of *Pelvetia canaliculata*, Orkney, and Clare Island.

BARYA Fuckel, *Symb. Mycol.* (1869), 93

Perithecia superficial, conoid or flask-shaped, separate, bright-coloured, soft, sometimes seated on a byssoid subiculum; asci elongated cylindrical or lanceolate, capitate; ascospores hyaline, linear, continuous, almost as long as the ascus.

Barya aurantiaca Plowr. & Wilson in *Gard. Chron.*, 9 February 1884, 176, figs. 32-4; *Claviceps Wilsoni* Cooke in *Grevillea*, xii, 77 (March 1884); *Claviceps purpurea* var. *Wilsoni* W. G. Sm., *Diseases of Field and Garden Crops* (1884), 233-8, figs. 107-11.

Subiculum white, floccose, becoming compact; perithecia almost superficial, conoid or flask-shaped, 0.25-0.3 mm. high, 0.12 mm. diameter, yellow, ostium orange; asci cylindrical, 200-250 \times 3 μ ; ascospores filiform, flexuous, continuous, almost as long as the ascus. Conidiophores spicarioid (?); conidia narrow-oval or lanceolate, ends acute, 10-12 \times 2-3 μ .

On *Claviceps purpurea* (Fr.) Tul. on ergot of *Glyceria fluitans*, near Aberdeen, July 1882 (A. S. Wilson), Hb. Kew.

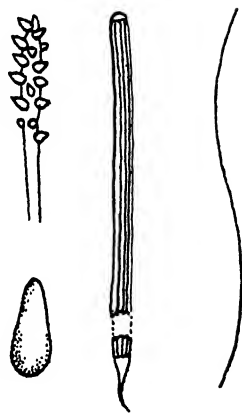


Fig. 24. *Barya aurantiaca*; perithecia on *Claviceps*, $\times 6$; a single perithecium, $\times 40$; ascus and spores, $\times 660$; ascospore, $\times 200$.

OPHIONECTRIA Sacc. in *Michelia* I (1878), 323

Perithecia superficial, separate, bright-coloured, sometimes clustered on a feebly developed, parenchymatous stroma; asci clavate, apex sometimes thickened; ascospores cylindrical or elongated fusoid, hyaline, multiseptate.

Ophionectria cylindrospora (Sollm.) Berl. & Vogl., *Add. Syll. Fung.* (1886), 217; *Nectria cylindrospora* Sollm. in *Bot. Zeit.* xxii (1864), 265.

Perithecia in small clusters on a feebly developed stroma within the bark, barely erumpent, globose, up to 0.3 mm. diameter, pale red, becoming dark red, blackening, encrusted with resinous particles and scales, collapsing and becoming cup-shaped; asci clavate or broadly clavate, 75-90 \times 8-20 μ , containing long cylindrical spores, or minute rod-like spores, or a mixture of the two; ascospores cylindrical, ends obtuse, or attenuated below, straight or curved, closely multiseptate, 22-40 \times 3-4 μ ; minute spores cylindrical, ends rounded, straight or curved, 2-4 \times 0.75 μ .



Fig. 25. *Ophionectria cylindrospora*; ascospore, $\times 500$.

On branches of *Pinus*. Lytchett, April 1922 (Hawley), Hb. B.M. On *Pinus Strobus*, Windsor Great Park, April 1935 (Ehrlich), Hb. Kew.

TRICHONECTRIA Kirschst. in *Verhandl. Bot. ver. Brand.* XLVIII (1906), 60

Perithecia superficial, solitary, or caespitose in small numbers, soft, bright-coloured, covered with long, spreading, hyaline setae; ascospores fusoid, hyaline, multiseptate.

Trichonectria hirta (Blox.) Petch in *Naturalist* (1937) 282; *Nectria hirta* Bloxam, in Currey, *Trans. Linn. Soc.* XXIV (1863), 158; *Calonectria hirta* (Blox.) Sacc. in *Michelia*, I (1878), 307; *Lasionectria hirta* (Blox.) Massee in *Grevillea*, xv (1886), 8; *Calonectria vermisporea* Mass. & Crossl. in *Naturalist*, January 1904, 4; *Dialonectria vermisporea* Mass. & Crossl., *Fungus Flora Yorks* (1905), 214.

Perithecia superficial, scattered or gregarious, minute, globose with a slightly prominent ostiole, or globoso-conoid, 0.1–0.25 mm. diameter, “pinkish salmon colour”, pallid when dry, covered with prominent white setae; setae hyaline, up to 110 μ long, 8 μ diameter below, tapering upwards, apex obtuse, slightly bulbous at the base, thick-walled below, solid above; asci soon diffluent, thin-walled, 14 μ diameter, apex rounded and not thickened; ascospores fusoid, straight or curved, or vermiform, ends obtuse, hyaline, multiseptate, 48–95 \times 5–9 μ .

On decaying fence rails, Twycross (Bloxam), Hb. Kew. and Hb. B.M. On a decorticated log, Hardcastle, near Hebden Bridge, Yorks (Crossland), Hb. Kew.

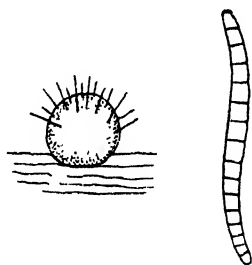


Fig. 26. *Trichonectria hirta*; perithecialium, $\times 40$; ascospore, $\times 350$.

TORRUBIELLA Boudier in *Rev. Mycol.* (1885), 227

Perithecia superficial, bright-coloured, membranous, sessile on a subiculum; asci linear, cylindrical, capitate; ascospores filiform, very long, septate, dividing into short part-spores, hyaline.

Torrubiella aranicida Boud., *loc. cit.*

Subiculum scanty, white; perithecia flask-shaped or elongated conoid, up to 0.7 mm. high, 0.25–0.35 mm. diameter, hyaline when fresh, becoming ochraceous or orange ochraceous when dry, straight or curved, glabrous; asci linear, very long, cylindrical, capitate, 250–350 \times 3–4 μ , four- or eight-spored; ascospores linear, very slender, almost as long as the ascus, 0.5–1 μ diameter, at first obscurely multiguttulate, then multiseptate with septa 8–16 μ apart, ultimately dividing into rod-shaped part-spores, 8–16 μ long.

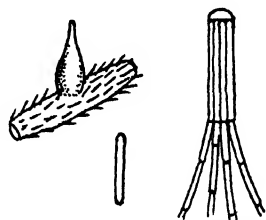


Fig. 27. *Torrubiella aranicida*; perithecialium on spider's leg, $\times 12$; apex of ascus and ascospores, $\times 1000$; part-spore, $\times 1000$.

On spiders under moss. Hubberholme, Yorks, 8 September 1936.

TRAILIA Sutherland in *Trans. Brit. Mycol. Soc.* v (1914), 149

Perithecia totally immersed, separate, pale coloured, soft, with a long beak; asci cylindrical, curved, eight-spored, without paraphyses; ascospores hyaline, linear clavate, bent double and coiled in the ascus, multiseptate.

Trailia Ascophylli Sutherland, *loc. cit.* pl. 3, fig. 2.

Perithecia gregarious, spherical or clavate, 50–60 μ diameter; beak cylindrical, straight or curved, 300–450 μ long, 7.5–10 μ diameter; asci 45–50 \times 9–10 μ ; ascospores narrow-clavate, tapering from 3.5 μ to 1 μ , hyaline, multiseptate, twice the length of the ascus (Sutherland).

In dark patches on *Ascophyllum nodosum*, Orkney.

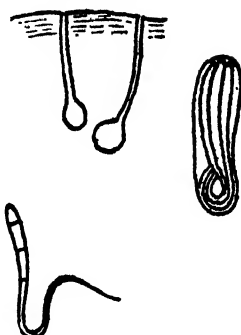


Fig. 28. *Trailia Ascophylli*; perithecia, ascus and ascospore (after Sutherland).

PLEONECTRIA Sacc., *Fungi Veneti nov. ser.* v, no. 178

Perithecia caespitose on a parenchymatous stroma or scattered, soft, bright-coloured; ascospores hyaline, muriform.

Pleonectria berolinensis Sacc. in *Michelia*, 1 (1878), 123; *Nectria Ribis* Rabenh., *Fung. Europ.* no. 247.

Perithecia erumpent, superficial, united in pulvinate groups, depressed globose, soon umbilicate-scutate, 0.3–0.5 mm. diameter, brick-red, ostium impressed; asci cylindrical, subsessile, 90–100 \times 10 μ , apex truncate, spores uniseriate; ascospores ovato-oblong, ends obtuse, muriform, with seven cross walls, not constricted, hyaline, 18–20 \times 8 μ (Saccardo, *loc. cit.*).



Fig. 29. *Pleonectria berolinensis*; ascospore, \times 750.

On dead branches of *Ribes* spp. Specimen in Hb. Kew., ex Hb. Cooke, sub *Nectria cinnabarina*, without name of host, date, or locality, but marked "British Fungi. M. C. Cooke"; det. Ehrlich.

HYPOCREACEAE

POLYSTIGMA DC. in *Comment. Mus. Hist.* III (1817), 330

Stroma fleshy, immersed in the leaf of the host, bright-coloured, circular or effused; perithecia immersed; ascospores hyaline, ovoid, continuous.

Polystigma rubrum (Pers.) DC., *loc. cit. supra*, 337; *Xyloma rubrum* Pers., *Observ.* II (1799), 101; Purton, *Midland Flora*, III (1821), 316; Greville, *Fl. Edin.* (1824), 365; Greville, *Scottish Crypt. Flora* (1824), pl. 120; *Dothidea rubra* Pers., Berkeley, *English Flora*, v (1836), 286.

Stromata generally scattered over the leaf, slightly thicker than the leaf, slightly arched, reddish yellow, fleshy, punctate with the ostiola of the pycnidia, subsequently (on the fallen leaves) strongly arched, concave above, almost black, brittle, brown internally; pycnidia immersed, spherical, chambered below, ostiolum small, papillate; pycnospores hyaline, linear, hook-shaped above, about 30μ long; perithecia immersed, almost spherical, ostiolum slightly projecting; asci oblongo-clavate, long-stalked, 78–87 (sporiferous part, 45–50) \times 10–12 μ , eight-spored; ascospores ellipsoid, continuous, hyaline, 11–13 \times 4.5 μ .

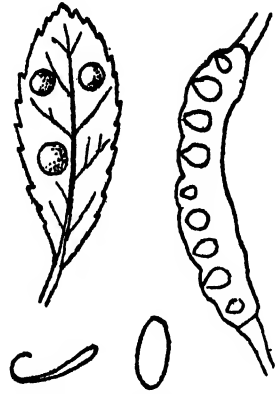


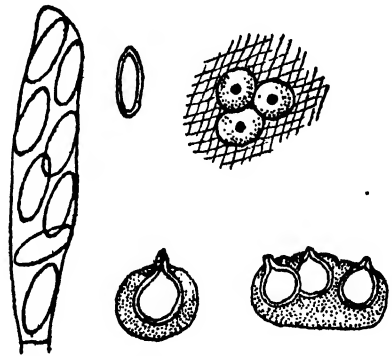
Fig. 30. *Polystigma rubrum*; affected leaf, natural size; cross-section of stroma, $\times 6$; pycnospore, $\times 450$; ascospore, $\times 660$ (after Tulasne).

On the leaves of *Prunus spinosa* and *Prunus insititia*. Common. Occurs also in Central Europe on *Prunus domestica*, and has been reported on that host from Thirsk, Yorks, in *Fungus Flora Yorks*.

Polystigma ochraceum (Wahlenb.) Sacc., *Conspectus Pyrenom.* (1875), 20; *Sphaeria ochracea* Wahlenb., *Flora Lapponica* (1812), 518; *Dothidea fulva* Holl. & Schm., Berkeley, *English Flora*, v (1836), 286.

Stromata circular or somewhat irregular, slightly arched, up to 1 cm. diameter, ochre-yellow, or golden yellow, becoming brownish red, closely punctate with the darker ostiola; in other respects, similar to *Polystigma rubrum*, but somewhat larger in all its parts; asci 95–105 \times 14 μ ; ascospores 14 \times 5–5.5 μ .

On leaves of *Prunus Padus*. "Not uncommon in Scotland" (Berkeley, *loc. cit.*): recorded for Dee and Moray in *Mycologia Scotica* (1879). "Highland Woods" (Klotzsch.), Hb. Kew.; without locality, Cooke, *Fungi Brit. Exsicc.* II, 578; near Aberdeen, 1869, Cooke, *Fungi Brit. Exsicc.* no. 564; Inellan, 1873 (R. H. Paterson), Hb. Kew.; Perth (J. B. White), Hb. Kew.



SELINIA Karsten in *Medd. Soc. Flora Fauna Fennica*, I (1876), 57

Stromata partly immersed, floccose, elliptic or wart-like, becoming confluent and irregular, bright-coloured, minutely tomentose; perithecia immersed; asci four- or eight-spored; ascospores ellipsoid, hyaline, continuous.

Selinia pulchra (Wint.) Sacc., *Syll. Fung.* II (1883), 457; *Hypocreopsis pulchra* Winter in *Hedwigia* (1875), 26.

Stromata at first small, subglobose or irregular, partly or almost

Fig. 31. *Selinia pulchra*; stroma seen from above, $\times 4$; simple stroma in section, $\times 10$; compound stroma in section, $\times 8$; ascus and spores, $\times 200$; ascospore, $\times 200$.

entirely immersed, containing a single perithecium, generally coalescing later into a rusty red crust, covered by a dense rusty brown tomentum, the hyphae of which bear short, cylindrical conidia; perithecia immersed, globose, up to 0.7 mm. diameter, with a stout, truncated conical, projecting ostium, wall yellow, ostium dark red; asci oblongo-clavate, sometimes ventricose below and attenuated above, sessile, apex rounded or truncate, $130-200 \times 40-48 \mu$, eight- or four-spored, spores biserial or irregularly clustered; ascospores ellipsoid, obtuse or subacute, hyaline, continuous, thick-walled, $48-55 \times 22-24 \mu$.

On sheep dung, Warleigh Common near Shrewsbury, October 1874 (Phillips and Plowright), Plowright, *Sphaer. Brit.* no. 100; on cow and sheep dung, Terrington St Clements, Norfolk, 1875 (Plowright); near King's Lynn, November-December, 1898 (Plowright).

HYPOCREOPSIS Karst., *Mycol. Fenn.* (1873), 251 and 281

Stroma superficial, effused, lobed, fleshy, bright-coloured; perithecia immersed; asci cylindrical, eight-spored; ascospores ellipsoid, hyaline, one-septate, not dividing.

Hypocreopsis lichenoides (Tode) Seaver in *Mycologia*, II (1910), 82; *Acrospermum lichenoides* Tode, *Fung. Meckl.* I (1790), 9; *Sphaeria riccioidea* Bolton, *Fung. Halifax*, IV (1791), 182; *Hypocrea riccioidea* (Bolt.) Berk., *Outlines* (1860), 383; *Hypocreopsis riccioidea* (Bolt.) Karst., *Myc. Fenn.* (1873), 251; Crossland in *Naturalist*, October 1908, 371, figs.

Stroma branched, the branches radiating from a common centre and much divided, with irregularly lobed ends, up to 10 cm. across, closely adherent to the substratum, pale fulvous brown or somewhat orange, darkening towards the centre, minutely tuberculate, fleshy, internally whitish, 4-6 mm. thick; perithecia immersed, subglobose; asci cylindrical, $170-180 \times 10 \mu$, eight-spored, spores obliquely uniserial; ascospores hyaline, oblong-fusiform, smooth, $24-30 \times 8-9 \mu$; paraphyses cylindrical, truncate above, about as long as the asci.

On dead branches of willow. Halifax (Bolton). Great Langdale, Westmoreland, May 1908 (Wheldon and Wilson), Hb. Kew. and Hb. B.M.; Corby Castle, Carlisle, 1883-6 (Dr Carlyle), Hb. Kew.; New Forest (Masse), Glamis, 1878 (Berkeley), Hb. Kew.; Dalry, Galloway, 9 September 1893 (J. McAndrew), Hb. B.M.

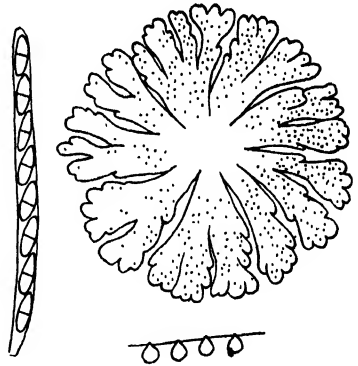


Fig. 32. *Hypocreopsis lichenoides*; stroma, $\times \frac{2}{3}$; perithecia (magnified); ascus and spores, $\times 200$ (after Crossland).

PODOSTROMA Karst. in *Hedwigia*, xxxi (1892), 294

Stroma stalked, clavate, erect, fleshy, light-coloured; perithecia immersed; asci cylindrical, eight-, then sixteen-spored; ascospores hyaline, at first ovoid, one-septate, then dividing into two globose, subglobose, or cuboid part-spores, hyaline or pale yellowish.

Podostroma alutaceum (Pers.) Atkinson in *Bot. Gaz.* XL (1905), 416; *Sphaeria alutacea* Pers., *Comment. Fung. Clavaef.* (1797), 12; *Sphaeria clavata* Sowerby, *Eng. Fungi* (1799), pl. 159; *Hypocrea alutacea* (Pers.) Tul., *Sel. Fung. Carp.* I (1861), 62; *Podocrea alutacea* (Pers.) Lindau, *Engler-Prantl, Naturl. Pflanzenf.* I, 1 (1897), 364.

Stroma vertical, clavate, up to 4 cm. high, sometimes with a rooting base, pale yellow or whitish, sterile below, fertile above; perithecia immersed, globose, 0.2–0.25 mm. diameter, with a papillate ostiolum; asci cylindrical, pedicellate, $80-90 \times 4 \mu$; part-spores globose, 4μ diameter, or oval, $4-4.5 \times 3 \mu$, hyaline, minutely warted.

On the ground in fir plantations, or on decaying stumps. Newmarket Heath (Rev. J. Hemsted); Costessy, Norfolk (Sowerby); Dinmore near Hereford, October 1874, Plowright, *Sphaer. Brit.* II, no. 3; Lucknam, Wilts, September 1843, 1859 (Broome), Hb. B.M.; Swanage, 19 November 1857 (Broome), Hb. B.M.; near King's Lynn (Plowright); Westley, Suffolk (Bloomfield); Haslemere (B.M.S.), Hb. Kew.; Den of Fullerton (Rev. J. Ferguson); etc.

HYPOCREA Fries, *Syst. Orb. Veg.* (1825), 104

Stroma horizontal, fleshy, pulvinate or effused, usually with a definite margin, bright-coloured; perithecia immersed; asci cylindrical, eight-, then sixteen-spored; ascospores at first ovoid, one-septate, then dividing into two globose, subglobose, or cuboid part-spores, hyaline or pale yellowish.

Hypocrea pulvinata Fuckel, *Symb. Mycol.* (1869), 185; *Hypocrea citrina* var. *fungicola* Karst., *Myc. Fenn.* II (1873), 204.

Stromata at first white and tomentose, becoming pale yellow and glabrous, punctate with the depressed ostiola, finally pale brown or pallid, circular or oval in plan, up to 8 mm. diameter, or larger and irregular by confluence, pulvinate, even, sometimes tuberculate when old, attached over the whole base, up to 2 mm. thick when fresh, margin definite rounded; context very pale yellow, almost white, becoming brownish when old, fleshy, becoming

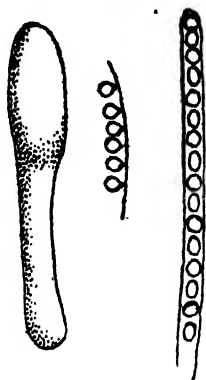


Fig. 33. *Podostroma alutaceum*; stroma, natural size; perithecia in section, $\times 6$; ascus and spores, $\times 500$.

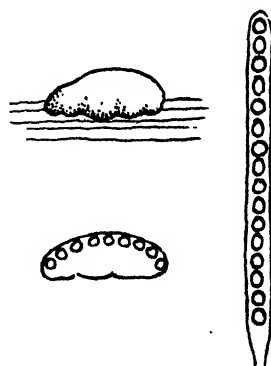


Fig. 34. *Hypocrea rufa*; stroma, $\times 6$; vertical section of stroma, $\times 6$; ascus and ascospores, $\times 500$.

corky; perithecia crowded in a yellow peripheral layer, small, ovoid, 0.2 mm. high, 0.15 mm. diameter, wall yellow, then brownish; asci cylindrical, $80-90 \times 3 \mu$; part-spores hyaline, smooth, broadly oval or rectangular, or somewhat wedge-shaped, $4-5 \times 2.5-3 \mu$, sometimes globose, 3μ diameter, extruded in white masses.

On decaying *Polyperi*, especially *Polyporus betulinus*. Darnaway Forest, September 1879 (Phillips and Plowright). Helmsley; North Wootton; Plumstead (Norfolk); Ludlow. Specimens in Hb. Kew. from Inver, Perth, 1911; Worcester, 25 May 1912 (J. W. Ellis); Mickleham, 11 November 1923 (J. Harvey Bloom); Black Hills, North Wootton, 30 March 1930 (T. Petch). Also recorded from Drumnadrochit, Aviemore, Minehead, Petersfield, Haslemere, and Arundel (B.M.S.).

Hypocrea splendens Phil. & Plowr. in *Grevillea*, XIII (1885), 79.

Stromata golden yellow, then reddish orange, with a brown tinge when old, dotted with brown ostiola, pulvinate or two-thirds globose, 3-6 mm. diameter, scattered, or clustered and distorted by mutual pressure, even, or minutely tuberculate when dry, base flat, internally fleshy, yellowish white; perithecia crowded in an orange peripheral zone, globose, 0.18 mm. diameter, or vertically oval, 0.25×0.15 mm., wall hyaline; asci cylindrical, $80-90 \times 4-5 \mu$, apex shortly capitate; part-spores cuboid, becoming globose, $4-5 \mu$ diameter, or oval, $4.5-6 \times 3.5-4 \mu$, hyaline, minutely warted.

On laurel sticks, Leicestershire, October 1881 (T. Howse), Hb. Kew.; Ken Wood, Hb. Kew., ex Hb. Cooke.

Hypocrea rufa (Pers.) Fr., *Summa Veg. Scand.* (1849), 383.

Stroma at first surrounded by a byssoid, white mycelium from which it separates, discoid, slightly convex, or pulvinate, 3-10 mm. diameter, sometimes confluent, at first chestnut-brown with a paler or white margin, becoming entirely chestnut-brown or rufous brown, opaque, faintly dotted with minute, subtranslucent ostiola, even, smooth, wrinkled when dry, fleshy, whitish internally, margin definite, rounded, sometimes incised or lobed, base flat; perithecia globose, about 0.2 mm. diameter, wall yellowish; asci cylindrical, sporiferous part, $60-75 \times 4-5 \mu$; part-spores globose, $3.5-4.5 \mu$ diameter, or oval, $5-6 \times 3.5-4 \mu$, hyaline, minutely warted.

Conidial stage, *Trichoderma lignorum* (Tode) Harz, effused, green, powdery; conidiophores lateral on the hyphae, solitary or in whorls, fusoid, 10μ high, 2μ diameter, bearing a small cluster of conidia on the attenuated apex; conidia spherical, blackish green, minutely warted, $3-6 \mu$ diameter.

On wood and bark; also on old fungi, *Polyperi*, *Pezizae*, *Elaphomyces*, etc. Generally distributed.

Hypocrea Schweinitzii (Fr.) Sacc., *Syll. Fung.* II (1883), 522; *Sphaeria Schweinitzii* Fr., *Elenchus*, II (1828), 60; *Sphaeria rigens* Fr., *Elenchus*, II (1828), 61; *Sphaeria contorta* Schwein. in *Trans. Amer. Phil.*

Soc. II (1832), 194; *Hypocrea contorta* (Schwein.) B. & C. in *Grevillea*, IV (1875), 14; *Hypocrea rigens* (Fr.) Sacc. in *Michelia*, I (1878), 301; *Hypocrea lenta* Auctt., non *Sphaeria lenta* Tode.

Stromata up to 1 cm. diameter, lenticular, centrally attached, circular, margin free and sometimes lobed, often undulating, black with a greenish tinge, minutely verrucose with projecting ostiola when mature, internally white, the isolated cells of the cortex green when fresh, olive to blackish when old; perithecia immersed, globose, about 0.15 mm. diameter, wall hyaline to pale brown; asci cylindrical, 60–66 \times 4 μ ; part-spores equal, globose, hyaline, almost smooth, with a large central gutta, 3.5 μ diameter.

On dead wood. Wood Heath, Chislehurst, August 1855 (Currey), Hb. Kew.; St Catherine's, November 1866 (Broome), Hb. Kew. and Hb. B.M.; Foxley Woods, October 1875 (Plowright and Phillips), Hb. B.M.; near Reeth, October 1936 (W. G. Bramley); Brandon, November 1876 (Plowright).

Hypocrea argillacea Phil. & Plowr. in *Grevillea*, XIII (1885), 79.

At first lemon-yellow, opaque, with slightly darker ostiola, about 2.5 mm. diameter, pulvinate, convex, even, glabrous, margin rounded, becoming dull brown, finally, or when dried, collapsing and becoming pale brownish yellow or red-brown, flattened pulvinate, or thin and discoid, minutely tuberculate with perithecial elevations; perithecia crowded in a peripheral layer, globose, 0.1–0.15 mm. diameter, or vertically ovoid, 0.15 mm. high, 0.1 mm. diameter, wall yellow; internally white, friable; asci cylindrical, 80–90 \times 3.5–4 μ ; part-spores unequal, the lower usually the larger, hyaline, minutely warted, globose, 3–3.5 μ diameter, or ovoid, 4–4.5 \times 3 μ .

The type specimen, on soft, rotten wood, Dersingham, November 1881, appears to have been lost. The above description has been drawn up from the following specimens. On a fallen branch of birch, North Wootton, November 1936; on dead wood, Lyndhurst (Massee) and Carlisle (Massee), both in Hb. B.M. as *Hypocrea farinosa*.

Hypocrea lutea Petch in *Journ. Bot.* LXXV (1937), 231; *Sphaeria gelatinosa* f. *lutea* Tode, *Fung. Meckl.* II (1791), 481.

At first white, circular, byssoid, plane, becoming fleshy, pulvinate or discoid, up to 2 mm. diameter, sometimes with a narrow byssoid margin, ochraceous when fresh (Hawley), red-brown when dry, sometimes subtranslucent above, minutely tuberculate with perithecial elevations; outer layer of the stroma and walls of the perithecia yellow-brown; internally white, friable, sometimes with a harder core at the base; perithecia globose or vertically ovoid; asci cylindrical, 90–100 \times 5 μ ; part-spores hyaline, minutely warted, unequal, globose, 4–4.5 μ diameter, or oval, 5 \times 3.5–4.5 μ , sometimes 7 \times 4 μ .

On dead leaves, Hurst Wood, October 1856 (Currey), Hb. Kew.; Tumby, Lincs, October 1905 (Hawley), Hb. B.M. The above description has been drawn up from the herbarium specimens. Currey at first considered his specimen a new species, but finally referred it to *Hypocrea gelatinosa*.

Hypocrea strobilina Phil. & Plowr. in *Grevillea*, xiii (1885), 79.

"Discoid, stroma whitish, thin, 1-4 mm. across; perithecia yellowish, rather large; asci cylindrical, octosporous; sporidia separating into two halves, each of which is subglobose, hyaline, $5-6 \times 5-5.5 \mu$."

On cones of spruce, Belmont, Hereford, November 1878 (J. Renny). Also recorded from Osmotherley, Yorks, on decaying pine wood (Crossland). No specimens available.

Hypocrea citrina (Pers.) Fr., *Summa Veg. Scand.* (1849), 383; *Sphaeria citrina* Pers., *Obs. Myc.* i (1796), 68.

Stroma effused, often spreading for several inches over wood, earth, etc., flat, often irregularly undulating or tuberculate owing to irregularities in the substratum, usually with a white, byssoid margin, elsewhere fleshy, lemon-yellow, minutely tomentose and minutely verrucose with the papillate, darker ostiola, internally white, solid, sometimes fibrillose at the base, up to 0.5 mm. thick; perithecia immersed, crowded in a yellow superficial zone, globose, 0.25 mm. diameter, wall yellow to brownish yellow; asci cylindrical, $85-100 \times 4-5 \mu$; part-spores unequal, the upper globose, 4μ diameter, the lower oval or subrectangular or wedge-shaped, $5-7 \times 3-4 \mu$, hyaline or pale yellow, minutely warted.

On dead wood, earth, etc. Generally distributed.

Hypocrea lactea Fr. is a white or pallid form of *H. citrina*, but the British specimens recorded under that name are misidentified.

Hypocrea placentula Grove in *Journ. Bot.* xxiii (1885), 133, pl. 256, fig. 5.

Stromata scattered, thin, at first with a byssoid margin, becoming compact with a definite rounded margin, discoid, suborbicular, 1-3 mm. diameter, flat, readily separable from the substratum, white, punctate with brownish ostiola; upper layer continuous, of crowded perithecia, internally loose; perithecia globose, 0.15 mm. diameter, almost completely immersed, wall pale yellow; asci cylindrical, $80-90 \times 3-4 \mu$; part-spores almost equal, globose, hyaline, minutely warted, $2.5-3.5 \mu$ diameter.

At the base of culms of *Juncus effusus*, Olton Reservoir, Warwickshire, September (Grove).

In some respects, the stroma resembles that of *Protocrea delicatula*, but the upper layer becomes continuous, and the perithecia do not project and are different in shape from those of the latter species.

CHROMOCREA Seaver in *Mycologia*, ii (1910), 58

Stroma, asci and spores as in *Hypocrea*, but spores coloured, green or olivaceous.

Chromocrea gelatinosa (Tode) Seaver, *loc. cit.*; *Sphaeria gelatinosa* Tode, *Fung. Meckl.* ii (1791), 48; *Hypocrea gelatinosa* (Tode) Fr.,

Summa Veg. Scand. (1849), 383; *Hypocrea moriformis* Cke & Massee, *Grevillea*, xvii (1888), 3.

Stromata pulvinate or hemispherical, 1–3 mm. diameter, often tuberculate, subgelatinous and subtranslucent when moist, the dark contents of the perithecia being visible, yellow or dark amber, becoming opaque and wrinkled when dry, internally pallid; perithecia immersed, crowded, vertically ovoid, up to 0.25 mm. high, 0.15 mm. diameter; asci cylindrical, $80-88 \times 3.5-4.5 \mu$, eight- then sixteen-spored; part-spores unequal, the upper globose, 4μ diameter, the lower oval or ellipsoid, $5-6 \times 3-4 \mu$, coarsely warted, dark green when fresh, becoming olivaceous when dry.

On decaying wood. Generally distributed.

Chromocrea cupularis (Fr.) Petch, comb. nov.; *Sphaeria cupularis* Fr., in *Linnaea* (1830), 539; *Hypocrea cupularis* (Fr.) Sacc., *Syll. Fung.* ii (1883), 535; *Hypocrea dacrymycella* Cooke & Plowr. in *Grevillea*, xii (1884), 100; *Hypocrea viscidula* Phil. & Plowr. in *Grevillea*, xiii (1885), 79.

Stromata at first globose, flattened above, then discoid, becoming cup-shaped, or ultimately pezizoid with an expanded or lobed margin, up to 10 mm. diameter, 5 mm. thick, pale yellow, externally viscid when fresh, but internally compact, whitish, becoming yellow; disc at first spotted with brown, then (when the spores are ripe) with green; perithecia confined to the disc, globose, 0.18 mm. diameter, or laterally oval, 0.18×0.12 mm., wall hyaline; asci cylindrical, capitate, $100-125 \times 7-8 \mu$; ascospores dividing equally or unequally; part-spores oval or subquadrangular, $8-9 \times 6-7 \mu$, or globose, $6-9 \mu$ diameter, with a few undivided (?) spores, oval, $12-13 \times 7 \mu$, dark green, becoming olivaceous, coarsely warted.

On coniferous logs. On *Pinus sylvestris*, Brandon, October 1881 (Plowright), Hb. Kew.

Chromocrea aureo-viridis (Plowr. & Cooke) Petch, comb. nov.; *Hypocrea aureo-viridis* Plowr. & Cooke in *Grevillea*, viii (1880), 104.

Stromata scattered or clustered, at first pulvinate, becoming depressed in the centre and pezizoid, centrally attached, circular, up to 2.5 mm. diameter, the larger examples plicate, pale yellow, then orange, the disc becoming olive, when dry pale brown with darker, subtranslucent ostiola, internally pale yellowish; perithecia confined to the disc, immersed, crowded, small, globose, about 0.1 mm. diameter; asci cylindrical, $70-75 \times 4 \mu$; part-spores equal, globose, 4μ diameter, or slightly ovoid, $4.5 \times 4 \mu$, dark green, coarsely warted.

On hazel. North Wootton, November 1879 (Plowright) Hb. Kew.

EPICHLÖE Fries, *Summa Veg. Scand.* (1849), 381 (as subgenus).

Stroma superficial, sessile, effused, surrounding the stems of grasses, at first white and byssoid, then fleshy, bright-coloured; perithecia at first free, then confluent; asci cylindrical, very long, capitate; ascospores linear, almost as long as the ascus, multiseptate, not dividing into part-spores.

Epichloë typhina (Pers.) Tul. in *Ann. Sci. Nat.* ser. 4, XIII (1860), 18; *Sphaeria typhina* Pers., *Icon. et Descript.* 1 (1798), 21; *Sphaeria spiculifera* Sow., *Engl. Fungi* (1803), pl. 274; *Dothidea typhina* (Pers.) Fr., *Syst. Mycol.* II (1823), 553; *Stromatosphaeria typhina* Grev., *Scott. Crypt. Flora*, IV (1826), pl. 204; *Hypocrea typhina* (Pers.) Berk., *Outlines* (1860), 383. Conidial stage, *Sphacelia typhina* (Pers.) Sacc. in *Michelia*, II (1881), 297.

Stroma sheathing the stems of grasses, up to 5 cm. long, at first white and bearing cylindrical or narrow-oval, hyaline conidia, $3-9 \times 1-2 \mu$, on *Cephalosporium* conidiophores, becoming orange or golden yellow; perithecia crowded, at first free, becoming confluent, oval, 0.3–0.6 mm. high, 0.25 mm. diameter, apex obtuse; asci $6-8 \mu$ diameter, eight-spored; ascospores linear, almost as long as the ascus, $1.5-2 \mu$ diameter, septate at intervals of $8-12 \mu$.

On living grass stems, especially *Dactylis*, *Holcus*, *Phleum*, etc. Common.

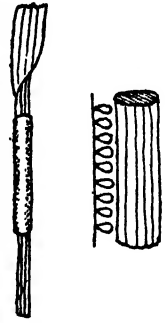


Fig. 35. *Epichloë typhina*; stroma on grass, $\times \frac{1}{2}$; stroma in longitudinal section $\times 6$.

OOMYCES B. & Br. in *Ann. Mag. Nat. Hist.* ser. 2, VII (1851), 185

Stroma erumpent, conoid, small, light-coloured, apex truncate, cortex membranous, tough, internally soft; perithecia few in each stroma, flask-shaped, vertical, opening on the truncate apex; asci narrow cylindrical, very long; ascospores linear, hyaline, multiseptate, not dividing into part-spores.

Oomyces carneo-albus (Libert) B. & Br., *loc. cit. supra*; *Sphaeria carneo-alba* Libert, *Crypt. Ard.* no. 241.

Stromata erumpent, scattered, pale flesh-coloured when fresh, becoming cream-coloured, matt, conical, 0.6–0.75 mm. high, 0.5 mm. diameter, contracted towards the apex, which is truncate, slightly convex, punctate with the mouths of the perithecia; perithecia three to seven in each stroma, immersed, narrow flask-

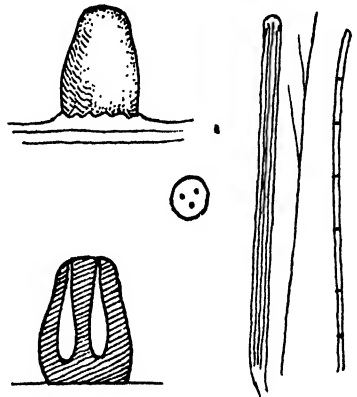


Fig. 36. *Oomyces carneo-albus*; stroma, $\times 20$; vertical section of stroma, showing perithecia, $\times 20$; apex of stroma, $\times 10$; ascus, paraphysis and ascospore, $\times 500$, but shortened.

shaped, vertical, up to 0.2 mm. diameter, almost as long as the height of the stroma, their apices fused with the wall of the stroma; asci 5–6 μ diameter, rather thick-walled, apex rounded, not capitate; paraphyses as long as the asci, 0.75 μ diameter, branched at the apex; ascospores linear, hyaline, multiseptate, 1.5 μ diameter, in a parallel bundle.

On dead leaves of *Aira caespitosa*. Spyre Park, Wilts (Broome), Hb. Kew. and Hb. B.M.; Lyndhurst (Masse), Hb. B.M.; Batheaston, January 1851, March 1859, Hb. Kew.; Carlisle, June 1886 (Dr Carlyle), Hb. Kew.; near Perth, May 1912 (J. Menzies), Hb. Kew.

CORDYCEPS Link, *Handb.* III (1833), 347

Stroma (clava) erect, usually consisting of a terete stalk and a clavate, ovoid, or subglobose head, fleshy; perithecia immersed in the head (in British species), usually flask-shaped, sometimes with the ostiola projecting; asci cylindrical, very long, capitate; ascospores filiform, almost as long as the ascus, multiseptate, dividing into short, rod-like, or narrow-oval, part-spores in the ascus.

Cordyceps ophioglossoides (Ehrh.) Link, *Handb.* III (1833), 347; *Sphaeria ophioglossoides* Ehrh., in Pers. *Holmsk. Coryph.* (1797), 144; *Sphaeria ophioglossoides* (Ehrh.) Fr., *Syst. Mycol.* II (1823), 324; *Clavaria parasitica* Willd., *Fl. Berol.* (1787), 405; *Cordyceps parasitica* (Willd.) Seaver in *North American Flora*, III (1910), 53.

Clava up to 10 cm. high, consisting of a comparatively slender stalk and an ovoid or oblong head very variable in size and shape, the whole fungus at first yellow or greenish yellow, becoming black; stalk smooth, 1–3 mm. diameter, terminating below in yellow mycelium; head usually laterally compressed, varying from narrow-oblong, 6 mm. high, 2 mm. broad, to broadly ovoid, 2.5 cm. high, 1.3 cm. broad, rounded above, smooth, then rough with projecting ostiola, becoming furrowed when old, internally loose and fuscous when dry; perithecia immersed, crowded, oval with a short neck, 0.7 mm. high, 0.36 mm. diameter; asci long, cylindrical, 7 μ diameter, eight-spored; part-spores hyaline, cylindrical with rounded ends, or oblong-oval, 2.5–5 \times 2 μ , a few subglobose, 2.5 μ diameter, with many longer, non-septate fragments, 6–30 \times 1.5–2 μ , often persisting in white masses on the mature fungus.

Parasitic on *Elaphomyces*. Generally distributed. Near Norwich (Berkeley, in *English Flora*, v, 233); Castle Rising (Plowright); North Wootton; Westwick.

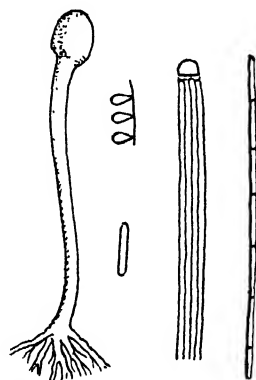


Fig. 37. *Cordyceps gracilis*; clava, natural size; perithecia in section, $\times 6$; upper part of ascus and ascospore, $\times 1000$; part-spore, $\times 1000$.

Recorded from Whitby, Doncaster, Minehead, Keswick, Drumnadrochit, Forres, Bettwsycoed (B.M.S.); and Thirsk, Raincliff Wood, Scarborough, Blackwood near Selby, Mulgrave Woods, Burnsall (Yorks. Nat. Union).

Cordyceps capitata (Holms.) Link, *Handb.* III (1833), 347; *Clavaria capitata* Holms., *Topsv.* 38 (1790); *Sphaeria capitata* (Holms.) Fr., *Syst. Mycol.* II (1823), 324; *Sphaeria agariciformia* Bolt., *Fung. Halifax* (1789), 130; *Cordyceps agariciformia* (Bolt.) Seaver in *North American Flora*, III (1910), 53.

Clavae solitary or clustered, up to 9 cm. high, consisting of a stout stalk and an ovoid or subglobose head, sharply defined from the stalk; stalk up to 7.5 cm. high, 1 cm. diameter, equal or slightly attenuated upwards, terete, smooth, becoming longitudinally furrowed, sometimes furfuraceous and hollow when old, yellow, blackening when old; head up to 2 cm. high and in diameter, yellow-brown or chestnut-brown, becoming black, smooth when fresh, rough with projecting ostiola when dry; perithecia immersed, oval or oblong-oval with a short neck, 0.6–0.7 mm. high, 0.25 mm. diameter; asci long, cylindrical, eight-spored, about 15μ diameter; part-spores hyaline, fusoid, narrow-oval, or subcylindrical, ends truncate and usually solid for a length up to 5μ , thick-walled, $7\text{--}48 \times 3\text{--}6\mu$.

Parasitic on *Elaphomyces*. Generally distributed, but not common. Ramsden Wood, Halifax (Bolton); Holt, Norfolk (Berkeley in *English Flora*, v, 233); Haslemere and Bettwsycoed (B.M.S.); West Runton and Westwick, Norfolk; Inverary (Stevenson and Paul); Weybridge; Botley, Hants (Phillips and Plowright); New Forest, — Costicles, Ramnor, Denny (Rayner).

Cordyceps militaris (Linn.) Link, *Handb.* III (1833), 347; *Clavaria militaris* Linn., *Sp. Pl.* (1753), 1182; *Torrubia militaris* Tul., *Sel. Fung. Carp.* III (1865), 6.

Clava up to 5 cm. high (usually smaller), 5 mm. diameter above, clavate, red or orange-red, waxy; head narrow-oval, apex rounded, slightly rough with the subtranslucent ostiola; stalk of the same colour as the head or rather paler; perithecia at first immersed, becoming more or less superficial in old and weathered, or dried, specimens, oval, up to 0.5 mm. high, 0.3 mm. diameter; asci cylindrical, capitate, very long, 4μ diameter, eight-spored; ascospores linear, as long as the ascus, dividing into short part-spores, which are at first cylindrical with truncate ends, becoming barrel-shaped or narrow-oval, $3.5\text{--}6 \times 1\text{--}1.5\mu$.

Conidial stage, *Cephalosporium* sp., occurring at the base of the perithecial clava and on the mycelium on the host; conidiophores simple, narrow flask-shaped or conical, up to 9μ high, 1.5μ diameter below, tapering to the apex; conidia hyaline, oval, or subpyriform with one end acute, $2.5\text{--}3 \times 1.5\text{--}2\mu$, sometimes globose, $1.5\text{--}2\mu$ diameter, with a few larger subpyriform with a truncate base, up to $6 \times 3\mu$.

On larvae and pupae of Lepidoptera. Generally distributed; usually occurring in the autumn.

Cordyceps gracilis Mont. & Dur. in *Flor. Alger*, 1, 449, pl. 25, fig. 2 (1846-69); *Cordiceps entomorrhiza* Berk. non (Dicks.) Link, *Outlines* (1860), 382, pl. 23, fig. 5, and Cooke, *Vegetable Wasps, etc.* (1892), 164; *Cordyceps Mawleyi* Westwood in *Gard. Chron.* ser. 3, ix (1891), 553, fig. 115.

Clavae up to 4 cm. high, consisting of a stout, usually straight stalk, up to 3.3 cm. high, and an ovoid or subglobose head, up to 7 mm. high, 5 mm. diameter; stalk terete, about 2 mm. diameter, yellow, smooth, expanding suddenly into the head; head chestnut-brown, smooth, ostiola inconspicuous; perithecia immersed, flask-shaped, 0.6 mm. high, 0.25 mm. diameter; asci cylindrical, capitate, very long, 4-5 μ diameter; ascospores linear, as long as the ascus, dividing into cylindrical, truncate, part-spores, 5-9 \times 1.5-2 μ .

Conidial stage, *Spicaria* (*Isaria*) *dubia* Delacr., in *Bull. Soc. Mycol. France*, ix (1893), 264, clavae numerous, slender, lax, decumbent or repent, up to 1 cm. long, 0.1 mm. diameter, white or yellowish, covered with diverging conidiophores; phialides in terminal or lateral clusters, flask-shaped, 7-10 \times 2-2.5 μ ; conidia hyaline, narrow fusoid, ends subacute, 4-6 \times 1 μ .

On buried larvae of Lepidoptera, usually *Hepialus*. Frequent; sometimes occurring in numbers in gardens in the spring. This species has usually been recorded in this country as *Cordyceps entomorrhiza*.

Cordyceps entomorrhiza (Dicks.) Link, *Handb.* III (1833), 347; *Sphaeria entomorrhiza* Dickson, *Plant. Crypt. Brit.* 1 (1785), 22, pl. 3, fig. 3; *Xylaria entomorrhiza* (Dicks.) Gray, *Nat. Arrang. Brit. Plants*, 1 (1821), 511.

Clava up to 19 cm. high, consisting of a slender, flexuose stalk and a subglobose head; stalk at first brown below, ashy above, becoming black, smooth, sometimes forked, about 1 mm. diameter; head subglobose or ovoid, about 6 mm. diameter, violet-grey, becoming black, rough with the projecting ostiola; part-spores cylindrical, truncate, 6-8 \times 1.5-2 μ (Tulasne).

On beetles and beetle larvae, usually *Carabus*. Found only once in this country, on a beetle larva, Bulstrode, Bucks, in the autumn (Dickson).

Cordyceps sphecocephala (Klotzsch) Cooke, *Vegetable Wasps, etc.* (1892), 40; *Sphaeria sphecocephala* Klotzsch in Hb. Hooker, Berkeley, *London Journ. Bot.* II (1843), 206.

Clava up to 6 cm. high, consisting of a slender stalk and a small ovoid or cylindrical head; stalk about 0.5 mm. diameter, usually flexuose and twisted, pale brown or brownish white, smooth, matt, longitudinally striate, the outer layer sometimes splitting away and recurving; head usually ovoid, acute above, up to 5 mm. high,

2–3 mm. diameter, sometimes cylindrical and up to 12 mm. long, yellow (? pale purple when fresh), dotted with subtranslucent ostiola, minutely vertically ridged when dry, with an ostiolum at the apex of each ridge; perithecia immersed, obliquely vertical, elongated flask-shaped or conoid, laterally compressed, 0.8 mm. high, 0.25 mm. broad; asci cylindrical, eight-spored, about $250 \times 8 \mu$; part-spores narrow oval, $8-15 \times 1.5-2.5 \mu$.

Conidial stage, *Hymenostilbe sphecophila* (Ditmar) Petch in *Trans. Brit. Mycol. Soc.* **xxi** (1937), 52; *Isaria sphecophila* Ditmar in *Sturm's Deutschl. Fl.*, Abt. III, Bd. I (1817), 115, pl. 57. Clava linear, up to 9 cm. high, 0.75 mm. diameter, tapering upwards, at first white or cream-coloured, then pale yellow or brownish yellow, straight or flexuose, terete, smooth, or very finely setulose in the upper part when magnified; basidia clavate or flask-shaped, $12-24 \times 4 \mu$; conidia clavate, upper end rounded, lower end truncate, hyaline, smooth, $6-10 \times 4-5 \mu$.

On Hymenoptera. ? On a wasp, Sluie, Rev. J. Keith, *Scottish Naturalist*, 1874; Specimen in Hb. B.M. ex Hb. Broome, from the Rev. G. Salmon, without locality or date. Conidial stage, Chopwell Wood, Northumberland, September 1933.

Cordyceps Forquignonii Quélet in *Compt. Rend. Assoc. Franc. Avanc. Sci.* (1887), 6, pl. 21, fig. 18; *Hypocrea myrmecophila* B. & Br. non Ces. in *Ann. Mag. Nat. Hist.* ser. 2, **vii** (1851), 186.

Clava up to 3.5 cm. high, consisting of a long, slender stalk and a small ovoid head; stalk irregularly curved, smooth, pale ochraceous, about 0.35 mm. diameter below, 0.5 mm. diameter above; head ovoid, 1.7 mm. long, 1 mm. diameter, orange; "asci cylindrical, $260-300 \times 5-6 \mu$; part-spores fusiform, $11-12 \times 2 \mu$ " (Rea in *Trans. Brit. Mycol. Soc.* **iv**, 314).

Conidial stage, *Hymenostilbe muscaria* Petch in *Naturalist*, April 1931, 101. Clava linear, terete, equal, or laterally compressed above and bifurcate at the apex, up to 3 cm. high, brownish white, matt, white pruinose above; basidia subcylindrical or clavate, $12-25 \times 2.5-3.5 \mu$, attenuated above, thick-walled, apex truncate or rounded, thickened, minutely verrucose, with one, or two, short, truncate sterigmata; conidia ovate, apex rounded, base obtuse or truncate, hyaline, smooth, $3-7 \times 1.5-3.5 \mu$.

On flies. Leigh Wood, Somerset, May 1846, 1847 (Broome), Hb. B.M.; Trefriw, North Wales, May 1874 (Phillips), Hb. B.M.; Higher Greenwood, Hebden Bridge, 6 May 1893 (N. & P.), Hb. Kew.; Masham, 8 August 1902 (W. A. Thwaites), Hb. Kew.; Grassington, 16 September 1931 (F. A. Mason); Dolgelly, B.M.S. Spring Foray, 1913; Bettwsycoed, B.M.S. Foray 1924; Ludlow, B.M.S. Spring Foray, 1932. Conidial stage, Arncliffe Woods, Whitby; Malyan Spout, Goathland; Holt House Wood, King's Lynn; Bolton Woods, Yorks.

OPHIOCORDYCEPS Petch in *Trans. Brit. Mycol. Soc.* xvi (1931), 73

Stroma (clava) as in *Cordyceps*, but asci clavate, thick-walled at the apex, and ascospores hyaline, elongated fusoid, multiseptate, not dividing into part-spores.

Ophiocordyceps clavulata (Schw.) Petch in *Trans. Brit. Mycol. Soc.* xviii (1933), 53; *Sphaeria clavulata* Schw., *Syn. N. A. F.* (1831), no. 1155; *Cordyceps clavulata* (Schw.) E. & E., *North American Pyreno.* (1892), 61, pl. 15; *Cordyceps pistillariaeformis* B. & Br. in *Ann. Mag. Nat. Hist. ser. 3*, vii (1861), 451, pl. 16, fig. 22.

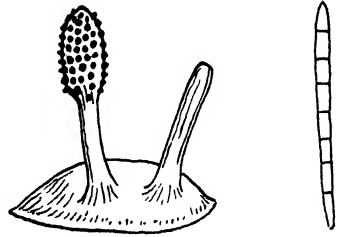


Fig. 38. *Ophiocordyceps clavulata*; scale insect bearing a perithecial and a conidial clava, $\times 10$; ascospore, $\times 500$.

Clavae gregarious, about 2 mm. high, consisting of a straight, terete stalk and a broader, ovoid or cylindrical head, the whole white, ashy, or pale brown, fibrillose; stalk equal or slightly attenuated upwards, 0.2–0.4 mm. diameter; head 0.5–1.2 mm. high, 0.4–0.6 mm. diameter, rough with the black apices of the perithecia; perithecia immersed, narrow-oval, conoid above, 0.25 mm. high, 0.12 mm. diameter, wall brown, apex black; asci narrow clavate, $80\text{--}100 \times 8\text{--}10 \mu$, eight-spored; ascospores narrow-clavate, seven- to eight-septate, $50\text{--}80 \times 2 \mu$.

Conidial stage, *Hirsutella lecanicola* (Jaap) Petch in *Trans. Brit. Mycol. Soc.* xviii (1933), 53; *Isaria lecanicola* Jaap in *Verh. Bot. Ver. Prov. Brandenburg* (1908), 49. Clavae white, ashy or pale brown, cylindrical or clavate, 2–4 mm. high, about 0.3 mm. diameter, apex obtuse; phialides crowded, base ovoid or conoid, $7\text{--}9 \times 4\text{--}5 \mu$, with a stout sterigma, $5\text{--}7 \mu$ long; conidia narrow-oval, oblong-oval, or clavate, $4\text{--}7 \times 1.5\text{--}2 \mu$.

On scale insects, *Lecanium* spp. On a scale insect on wych elm, Batheaston, Somerset, March 1860, October 1860 (Broome), Hb. Kew. and Hb. B.M.

CLAVICEPS Tulasne in *Compt. Rend. Acad. Sci. Paris*, xxxiii (1851), 646

Stroma (clava) erect, arising from a sclerotium (ergot) and consisting of a thin, terete stalk and a globose head, bright-coloured, soft; perithecia immersed in the head, usually flask-shaped, sometimes with the ostiola projecting; asci cylindrical, very long, capitate; ascospores hyaline, linear, continuous in the ascus, but becoming septate on germination after extrusion.

Claviceps purpurea (Fr.) Tul. in *Ann. Sci. Nat. ser. 3*, xx (1853), 45, pls. 1, 2, 3, figs. 1–11; *Sphaeria purpurea* Fr., *Syst. Mycol.* ii (1823), 325; *Cordiceps purpurea* Fr., Berkeley, *Outlines* (1860), 382; *Claviceps microcephala* (Wallr.) Tul. in *Ann. Sci. Nat. ser. 3*, xx (1853), 49;

Kentrosporium microcephalum Wallr. in *Beitr. Bot.* 1 (1844), 164; *Cordiceps microcephala* Tul., Berkeley, *Outlines* (1860), 382; ? *Sphaeria Hookeri* Kl., in Berkeley, *English Flora*, v (1836), 234. Conidial stage, *Sphacelia segetum* Lév. in *Mém. Soc. Linn.* v (1827), 578; *Farinaria Poae* Sow. pl. 396, f. 6. Ergot, *Spermoedia clavus* (DC.) Fr., *Syst. Mycol.* II (1822), 268; *Sclerotium Clavus* DC., *Fl. Franc.* VI (1815), 115.

Conidial stage, white covering the ovary and adjacent structures; conidia ellipsoid, hyaline, $4-6 \times 2-3 \mu$. Ergot, black, subcylindrical or fusoid, usually angular in section, straight or curved, sometimes longitudinally grooved, up to 2 cm. long, formed beneath the ovary, and pushing off the remains of the latter and the conidial stage. Perithecial clavae arising from the fallen ergot, and consisting of a long or short stalk and a subglobose head; stalk slender, straight or flexuose, purple, subtranslucent when fresh, smooth, socketed into the head; head depressed globose, cream-coloured, with brown ostiola, opaque, or purple and subtranslucent, even or tuberculate, the differences depending upon humidity during growth; perithecia immersed, flask-shaped, ostiola projecting slightly when dry; asci cylindrical, capitate, very long, about 5μ diameter, eight-spored; ascospores long, linear, $0.75-1 \mu$ diameter, continuous.

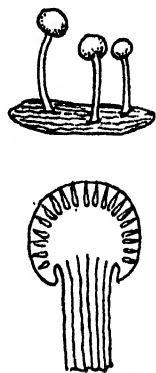


Fig. 39. *Claviceps purpurea*; clavae on ergot, $\times 2$; longitudinal section of the head, showing perithecia, $\times 8$.

On the inflorescence of various grasses. Common. Several biologic forms occur.

Claviceps nigricans Tul. in *Ann. Sci. Nat.* ser. 3, xx (1853), 51, pl. 4, figs. 15-22.

Ergot linear, semicylindrical, obtuse, often strongly curved, black, up to 12 mm. long. Perithecial clava up to 8 mm. high, entirely blackish violet, head at first paler; head depressed globose, minutely verrucose; perithecia flask-shaped, ostiola slightly projecting; asci very long, cylindrical, capitate, eight-spored; ascospores linear, continuous, hyaline.

On *Scirpus* spp. Only the ergot has been observed in this country. On *Heleocharis palustris*, Hengistbury Head, and on ? *Heleocharis uniglumis*, Poole Heath, Dorset, August 1863 (H. Trimen), Hb. Kew.; on *Heleocharis palustris*, Clatto Reservoir, Fife, October 1909 (G. West), Hb. Kew.

REJECTED NAMES

Broomella leptogicola Cke & Massee is the fructification of a lichen.

Broomella Vitalbae B. & Br. belongs to the Sphaeriaceae.

Calonectria Leightonii (B. & Br.) Sacc. (*Nectria Leightonii* B. & Br.) is a lichen.

Melanospora discospora Massee & Salmon is probably *Chaetomium murorum* Cda.

Melanospora gigantea Massee & Crossland is *nomen nudum*.

Melanospora vitrea (Cda) Sacc. is *Sphaeronaemella*.

Nectria caulina Cke is *Ditopella Vizeana* Sacc. & Speg. (Sphaeriaceae).

Nectria umbrina (Berk.) Fr. belongs to the Sphaeriaceae.

SPECIES DESCRIBED AS NEW IN THIS PAPER

Calonectria tessellata Petch, n.sp.

Peritheciis superficialibus, sparsis vel gregariis, udo sordide auran-
tiacis, sicco brunneo-flavis, conicis vel ovoideis, ad 0.3 mm. alt.,
0.25 mm. diam., interdum medio collapsis, minute rugosis, glabris
vel apice leniter verrucosis, ostiolo nudo, obscuriori; pariete externe
cellularum magnarum composito; ascis clavatis, octosporis, sessilibus,
apice truncato, $75-95 \times 12-18 \mu$, sporis oblique uniseriatis vel biseria-
tis; ascosporis hyalinis, triseptatis, ovalibus vel oblongo-ovalibus,
interdum inaequilateralibus, rectis, obtusis, $18-26 \times 7-9 \mu$.

On decaying stalks of *Brassica*, North Wootton, Norfolk, November 1935; on
dead apple twig, Camberley, November 1920, Hb. B.M.

Gliocladium strictum Petch, n.sp.

Conidiophoris congestis, albis, circa 100μ alt., stipite basi 4μ diam.,
ramulis paucis; ramulis inferioribus saepius solitariis, distantibus,
superioribus (phialidibus) oppositis vel tri-verticillatis, circa 30μ
longis, attenuatis, ramulis omnibus erectis, parallelis; conidiis hya-
linis, oblongo-ovalibus vel angusto-ovalibus, leniter inaequilatera-
libus, obtusis, $5-11 \times 2-2.5 \mu$, longioribus ultimo uniseptatis vel
pseudo-uniseptatis, muco conglutinatis.

On *Fomes annosus*; the conidial stage of *Hypomyces Broomeanus*.

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LIST OF MEMBERS

Correct to 12th May, 1938.

Honorary Members

- Lister, Miss Gulielma, F.L.S., 871 High Road, Leytonstone, Essex. (1903.) (1924.)
 Rea, Mr Carleton, B.C.L., M.A., 6 Barbourne Terrace, Worcester. (1896.) (1918.)

Ordinary Members

1. Aberdeen, The University Library. (1916.)
2. Adams, Rev. J. H., Landulph Rectory, Hatt, Saltash, Cornwall. (1919.)
3. Adcock, Mr Archie, Upton Road, Norwich. (1921.)
4. Ainsworth, Mr G. C., B.Sc., Ph.D., Experimental and Research Station, Cheshunt, Herts. (1931.)
5. Alaily, Mr Y. A. S. El, The Botany School, Cambridge. (1935.)
6. Alberta, University of Edmonton, Alberta, Canada. (1924.)
7. Alcock, Mrs N. L., F.L.S., M.B.E., 12 Tavistock Square, London, W. 1. (1919.)
8. Ashby, Mr S. F., B.Sc., Imperial Mycological Institute, Ferry Lane, Kew, Surrey. (1926.)
9. Bacon, Mrs. Alice, B.Sc., F.L.S., Technical College, Brighton, Sussex. (1938.)
10. Barnes, Mr B., D.Sc., Ph.D., V.-P.L.S., Chelsea Polytechnic, London, S.W. 3. (1922.)
11. Barr, Rev. Robert, T.D., M.A., The Manse, Neilston, Renfrewshire. (1918.)
12. Barrington, Dr F. J. F., University College Hospital, Medical School, University Street, London, W.C. 1. (1901.)
13. Bartlett, Mr A. W., M.A., M.Sc., F.L.S., Department of Botany, King's College, Newcastle-on-Tyne. (1920.)
14. Bates, Mr G. R., c/o British South Africa Company, Mazoe Citrus Estate, Mazoe, S. Rhodesia. (1930.)
15. Bates, Mrs L. F., B.Sc.,
16. Beardslee, Mr H. C., Perry, Ohio, U.S.A. (1933.)
17. Beaumont, Mr Albert, M.A., Seale-Hayne Agricultural College, Newton Abbot, Devon. (1924.)
18. Bewley, Mr W. F., D.Sc., Experimental and Research Station, Cheshunt, Herts. (1922.)
19. Biffen, Professor Sir Rowland H., M.A., F.R.S., 136 Huntingdon Road, Cambridge. (1899.)

20. Biggs, Miss R., B.Sc., Ph.D., 26 *b* Newton Road, London, W. 2.
(1937.)
21. Biologist, Plant Research Laboratory, Horticultural Gardens,
Burnley, Victoria, Australia. (1921.)
22. Birmingham Natural History and Philosophical Society, c/o
Mrs O. W. Thompson, 18 Hermitage Road, Edgbaston,
Birmingham. (1920.)
23. Bisby, Mr Guy R., Ph.D., Imperial Mycological Institute,
Ferry Lane, Kew, Surrey.
24. Blackman, Professor V. H., M.A., F.R.S., F.L.S., 17 Berkeley
Place, Wimbledon, London, W. 19. (1900.)
25. Blackwell, Miss E. M., M.Sc., F.L.S., Botanical Department,
Royal Holloway College, Englefield Green, Surrey. (1917.)
26. Blumer, Dr S., Myrtenweg 12, Bern-Bumpliz, Switzerland.
(1936.)
27. Bonn, Germany, Institut für Pflanzenkrankheiten, Nuss-Allee
9. (1931.)
28. Boston, The Mycological Club, Horticultural Hall, Boston,
Mass, U.S.A. (1926.)
29. Bourgin, Dr Viennot, École Nationale d'Agriculture de
Grignon, Seine-et-Oise, France. (1936.)
30. Braid, Professor K. W., B.A., B.Sc., West of Scotland Agri-
cultural College, 6 Blythswood Square, Glasgow. (1922.)
31. Brazier, Mr E., Ty'n-y-gongl, Love Lane, Stourbridge. (1921.)
32. Brenchley, Mr G. H., B.A., Clare College, Cambridge. (1925.)
33. Brett, Miss M., M.Sc., Ph.D., Northern Polytechnic, Holloway
Road, London, N. 7. (1921.)
34. Brierley, Professor W. B., D.Sc., F.R.A.I., F.L.S., Department
of Agricultural Botany, The University, Reading. (1919.)
35. Brinton, Mr R. E. B., 68 Woodstock Avenue, Golders Green,
London, N.W. 11. (1935.)
36. Brisbane, The Director, Bureau of Sugar Experiment Stations,
Department of Agriculture and Stock, Queensland,
Australia. (1930.)
37. British Museum, The Trustees of, Cromwell Road, South
Kensington, London, S.W. 7. (1914.)
38. Brooks, Professor F. T., M.A., F.R.S., F.L.S., The Botany
School, Cambridge. (1907.)
39. Brown, Dr Mabel Raven, Newnham College, Cambridge. (1936.)
40. Brown University, Library, East side Station, Providence, R.I.,
U.S.A. (1920.)
41. Brown, Professor W., M.A., D.Sc., F.R.S., Imperial College of
Science, South Kensington, London, S.W. 7. (1922.)
42. Bruxelles, Jardin Botanique de l'État, c/o M. P. van Aerdschot.
(1911.)

43. Buckley, Mr W. D., "St Anthony", Leigh Park, Datchet, Bucks. (1916.)
44. Buddin, Mr Walter, M.A., Laboratory of Plant Pathology, University of Reading, 7 Redlands Road, Reading. (1921.)
45. Buller, Professor A. H. R., D.Sc., Ph.D., F.R.S., c/o Herbarium, Royal Botanic Gardens, Kew. (1911.)
46. Bunting, Mr R. H., F.L.S., 3 Stanton Court, Weymouth. (1921.)
47. Burges, Mr N. A., The Botany School, Cambridge. (1935.)
48. Burr, Mr S., M.Sc., Department of Agriculture, The University, Leeds. (1924.)
49. Butler, Mr E. J., C.I.E., C.M.G., D.Sc., M.B., F.R.S., F.L.S., Agricultural Research Council, 6A Dean's Yard, London, S.W. 1. (1920.)
50. Caldwell, Mr J., D.Sc., Ph.D., Department of Botany, University College, Exeter. (1932.)
51. Cambridge, The Botany School. (1920.)
52. Campbell, Mr A. H., B.Sc., Ph.D., Department of Botany, The University, Bristol. (1934.)
53. Carne, Mr W. M., F.L.S., The University of Tasmania, Hobart, Tasmania, Australia. (1928.)
54. Carr, Professor J. W., M.A., F.L.S., Mapperley Edge, Private Road, Sherwood, Nottingham. (1896.)
55. Carrothers, Mr E. N., 7 Fitzwilliam Street, Belfast, N. Ireland. (1925.)
56. Cartwright, Mr K. St G., M.A., F.L.S., The Red House, Kingston Blount, Oxford. (1913.)
57. Castellani, Sir Aldo, M.D., 23 Harley Street, London, W. 1. (1922.)
58. Cayley, Miss Dorothy M., John Innes Horticultural Institution, Mostyn Road, Merton Park, London, S.W. 19. (1913.)
59. Charles, Miss Vera K., United States Department of Agriculture, Bureau of Plant Industry, Washington D.C., U.S.A. (1933.)
60. Chaudhuri, Mr H., M.Sc., Ph.D., University of the Punjab, Lahore, India. (1920.)
61. Cheal, Mr W. F., Savile House, Queen's Road, Wisbech, Cambs. (1927.)
62. Chesters, Mr C. G. C., B.Sc., M.Sc., Ph.D., Botanical Department, The University, Edgbaston, Birmingham. (1930.)
63. Ciferri, Professor Dr R., Assistant Director, Laboratorio Crittogamico, Casella Postale 165, Pavia, Italy. (1926.)
64. Clapham, Mr A. R., M.A., Ph.D., Department of Botany, The University, Oxford. (1931.)
65. Cleland, Mr J. Burton, M.D., Professor of Pathology, University of Adelaide, South Australia. (1918.)

66. Clouston, Mr D., M.A., B.Sc., (Agr.), North of Scotland College of Agriculture, Crown Mansions, 41 Union Street (2nd Floor), Aberdeen. (1931.)
67. Colson, Miss B., B.Sc., Ph.D., The University, Manchester. (1934.)
68. Connecticut Agricultural Experiment Station, New Haven, Connecticut, U.S.A. (1929.)
69. Cook, Mr W. R. I., B.Sc., Ph.D., Department of Botany, University College, Newport Road, Cardiff. (1924.)
70. Cooke, Mr G. J., 143 Newmarket Road, Norwich. (1933.)
71. Cooke, Mrs G. J., 143 Newmarket Road, Norwich. (1937.)
72. Cooper, Miss Charlotte A., California Lane, Bushey Heath, Herts. (1911.)
73. Cooper, Mrs V. Astley, The Tors, Knowle, Fareham, Hants. (1921.)
74. Cornell University, The Library, New York State College of Agriculture, Ithaca, N.Y., U.S.A. (1920.)
75. Corner, Mr E. J. H., M.A., F.L.S., Assistant Director, Botanic Gardens, Singapore, Straits Settlements. (1924.)
76. Cotton, Mr Arthur D., O.B.E., F.L.S., Keeper, Herbarium, Royal Botanic Gardens, Kew, Surrey. (1902.)
77. Croxall, Mr H. E., B.Sc., Research Station, Long Ashton, Bristol. (1937.)
78. Cunningham, Mr G. H., Ph.D., Plant Research Station, Box 442, Palmerston North, New Zealand. (1922.)
79. Curtis, Miss Kathleen M., M.A., D.Sc., D.I.C., F.L.S., Mycologist, Biological Department, Cawthron Institute of Scientific Research, Nelson, New Zealand. (1917.)
80. Cutting, Mr E. M., M.A., F.L.S., Botanical Department, University College, Gower Street, London, W.C. 1. (1920.)
81. Dade, Mr H. A., A.R.C.S., Imperial Mycological Institute, Ferry Lane, Kew, Surrey. (1927.)
82. Das, Mr Kedarnath, C.I.E., M.D., Principal, Carmichael Medical College, 1 Belgachia Road, Calcutta, India. (1922.)
83. Davies, Mr D. W., B.Sc., Adviser in Mycology, Agricultural Buildings, University College of Wales, Aberystwyth. (1923.)
84. Davies, Mr D. L. Godfrey, Research Station, Long Ashton, Bristol. (1938.)
85. Day, Mr W. R., B.A., B.Sc., Imperial Forestry Institute, Oxford. (1928.)
86. Deacon, Dr G. E., Brundall, Norwich. (1933.)
87. Dehra Dun, The Forest Botanist, Forest Research Institute and College, U.P., India. (1929.)

88. Deighton, Mr F. C., M.A., Mycologist, Department of Lands and Forests, Freetown, Sierra Leone, West Africa. (1925.)
89. Delhi, Imperial Mycologist, Imperial Agricultural Research Institute, Delhi, India. (1921.)
90. Dennis, Mr R. W. G., Ph.D., School of Agriculture, Cambridge. (1932.)
91. Dickinson, Mr S., Ph.D., School of Agriculture, Cambridge. (1921.)
92. Dobbs, Mr C. G., B.Sc., Ph.D., Botanical Department, King's College, Strand, London, W.C. 2. (1933.)
93. Dodge, Dr Carroll W., Missouri Botanical Garden, 2315 Tower Grove Avenue, St Louis, Missouri, U.S.A. (1926.)
94. Dowson, Mr W. J., M.A., D.Sc., The Botany School, Cambridge (1920.)
95. Duff, Mr R. G., Perly Cross, Teignmouth, Devon. (1938.)
96. Duncan, Mr J. T., London School of Hygiene and Tropical Medicine, Keppel Street, London, W.C. 1. (1930.)
97. Dunston, Capt. Ambrose E. A., Burltons, Donhead St. Mary, Wiltshire (via Shaftesbury). (1937.)
98. Edwards, Mr W. H., Belle Vue, Barline, Beer, Devon. (1896.)
99. Elliott, Mrs J. S. Bayliss, D.Sc. (B'ham.), B.Sc., (London), Arden Grange, Tanworth-in-Arden, Warwickshire. (1911)
100. Ellis, Mr E. A., Castle Museum, Norwich. (1937.)
101. Ellis, Mr E. H., B.Sc., Gramarye, Farley Green, Guildford, Surrey. (1936.)
102. Ellis, Miss E. M., B.A., B.Sc., St Hugh's College, Oxford. (1930.)
103. Ellis, Mr Holmes, F.R.M.S., 108 Birtwistle Avenue, Colne, Lancs. (1927.)
104. Emerson, Mr R., The Botany School, Cambridge. (1938.)
105. Emoto, Dr Y., Biological Department, Peers' College (Gaku-shuin), Mejiromachi, Tokyo, Japan. (1929.)
106. Essex Field Club, c/o Mr Percy Thompson, F.L.S., Essex Museum of Natural History, Romford Road, Stratford, London, E. 15. (1919.)
107. Exeter, Librarian, University College of the South-West of England. (1926.)
108. Eyre, Miss J. C., c/o Miss Wakefield, Herbarium, Royal Botanic Gardens, Kew. (1915.)
109. Fenton, Mr E. W., M.A., B.Sc., F.L.S., Biology Department, Edinburgh and East of Scotland College of Agriculture, Edinburgh. (1920.)
110. Findlay, Mr W. P., B.Sc., A.R.C.S., Courte Falaise, Sevenoaks, Kent. (1928.)
111. Finlayson, Mr Raymond A., F.L.S., Official Seed Testing Station, Huntingdon Road, Cambridge. (1910.)

- 112. Fisher, Mr S. D. P., Sackville Street, Leeds. (1930.)
- 113. Fitzpatrick, Professor H. M., Ph.D., 220 Bryant Avenue, Ithaca, New York, U.S.A. (1935.)
- 114. Fountain, Mr A. S., F.R.M.S., "Thornham" Fulwood Road, Sheffield, Yorks. (1934.)
- 115. Fraser, Miss Lilian R., D.Sc., Bank of New South Wales, Threadneedle Street, London. (1938.)
- 116. Gadd, Mr C. H., D.Sc., Tea Research Institute, Nuwara Eliya, Ceylon. (1921.)
- 117. Galloway, Mr L. D., 31 Lake Close, Wimbledon, London, S.W. 19. (1928.)
- 118. Gardner, Capt. Frederic, c/o Barclays Bank, Jersey, C.I. (1898.)
- 119. Garrett, Mr S. D., Rothamsted Experimental Station, Harpenden, Herts. (1936.)
- 120. Garside, Mr S., M.Sc., F.L.S., Botanical Department, Bedford College, Regent's Park, London, N.W. 1. (1922.)
- 121. Gates, Professor R. R., D.Sc., Ph.D., F.R.S., F.L.S., King's College Strand, London, W.C. 2. (1921.)
- 122. Ghamrawy, Mr Ali K., 39 Monirah Street, Cairo, Egypt. (1932.)
- 123. Gibson, Miss C. J., B.A., 27 Banbury Road, Oxford. (1933.)
- 124. Gilbert, Dr E. M., Botanical Department, University of Wisconsin, Madison, Wis., U.S.A. (1922.)
- 125. Gilbert, M. E., Docteur en Pharmacie, 6 Rue de Laos, Paris (15^e), France. (1924.)
- 126. Gill, Mr G. E., LL.B., 68 Pembroke Road, Dublin, Irish Free State. (1937.)
- 127. Gillespie, Mr J., B.Sc., Botany Dept., University of Reading, (1938.)
- 128. Glasstone, Mrs V. F. C., B.A. (Oxon.), 15 Northumberland Road, Sheffield. (1930.)
- 129. Glynne, Miss Mary D., M.Sc., F.L.S., Rothamsted Experimental Station, Harpenden, Herts. (1932.)
- 130. Gorman, Mr M. J., A.C.C.Sc.I., Albert Agricultural College, Glasnevin, Dublin. (1925.)
- 131. Gould, Mr F. G., Woodrising, Trapps Hill, Loughton, Essex. (1918.)
- 132. Green, Col. C. Theodore, A.M.S., M.R.C.S., L.R.C.P., F.L.S., 31 Shrewsbury Road, Birkenhead. (1901.)
- 133. Green, Miss F. Mary, M.A., Ph.D., 63 Canford Road, Clapham Common, London, S.W. 11. (1931.)
- 134. Gregor, Mrs M. J. F., Ph.D., Royal Botanic Garden, Edinburgh. (1927.)
- 135. Gregory, Mr P. H., Ph.D., Seale Hayne Agricultural College, Newton Abbot, Devon. (1930.)

136. Grieve, Mr B. J., M.Sc., Botanical Department, Melbourne University, Carlton, Victoria, Australia. (1931.)
137. Grinling, Mr C. H., B.A., 71 Rectory Place, Woolwich. (1913.)
138. Gwynne-Vaughan, Professor Dame Helen, G.B.E., D.Sc., LL.D., F.L.S., 93 Bedford Court Mansions, London, W.C. 1. (1906.)
139. Halley, Miss E. M., The Botany School, Cambridge. (1937.)
140. Hanna, Mr W. F., M.Sc., Dominion Rust Research Laboratory, Agricultural College, Winnipeg, Canada. (1925.)
141. Hansford, Mr C. G., M.A., F.L.S., Mycologist, Department of Agriculture, Kampala, Uganda. (1921.)
142. Harley, Mr J. L., M.A., D.Phil., Shrublands, Hethersett, Norfolk. (1932.)
143. Harris, Mr G. C. M., 148 Divinity Road, Oxford. (1934.)
144. Harris, Mr R. V., B.Sc., A.R.C.S., Horticultural Research Station, East Malling, Kent. (1924.)
145. Harrison, Mr T. H., D.Sc., Australia House, Strand, London, W.C. 2. (1931.)
146. Harvard University, The Library, Cambridge, Mass., U.S.A. (1923.)
147. Hastings, Mr Somerville, M.S., F.R.C.S., 43 Devonshire Street, Portland Place, London, W. 1. (1913.)
148. Hawker, Miss L. E., Ph.D., Botanical Department, Imperial College of Science, London, S.W. 7. (1934.)
149. Heim, M. Roger, Sous-Directeur au Muséum d'Histoire Naturelle, 11 Rue de Médicis, Paris (6^e), France. (1930.)
150. Heimbeck, Mrs Louise, Brosoe, Levanger, Norway. (1923.)
151. Hemmi, Dr Takewo, Phytopathological Institute, Department of Agriculture, Kyoto Imperial University, Kyoto, Japan. (1923.)
152. Hereford, Mr E. H., 131 Queen Victoria Street, London, E.C. 4 (1933.)
153. Hickman, Mr C. J., c/o Plant Pathology Lab., Milton Rd., Harpenden, Herts. (1935.)
154. Hildyard, Mr F. W., 1 Lichfield Road, Kew, Surrey. (1913.)
155. Holden, Professor H. S., D.Sc., F.R.S.E., F.L.S., Forensic Science Laboratory, Burton Street, Nottingham. (1923.)
156. Honolulu, Association of Hawaiian Pineapple Cannerys, P.O. Box 3166, Hawaii. (1929.)
157. Honolulu, The Library, Experimental Station, S.P.A., Box 411, Hawaii. (1920.)
158. Hopkins, Mr J. C. F., D.Sc., A.I.C.T.A., P.B., 74B Salisbury, S. Rhodesia. (1930.)
159. Horne, Mr A. S., D.Sc., F.L.S., F.G.S., Botanical Department, Imperial College of Science, South Kensington, London, S.W. 7. (1921.)

160. Howard, Mr H. J., F.R.M.S., F.L.S., "Lingfield", 6 College Road, Norwich. (1918.)
161. Hubbard, Miss M. D., B.Sc., Laren, Western Drive, Littleover, Derby. (1933.)
162. Hughes, Mr G. C., Priory Road, Bicester. (1898.)
163. Hughes, Mr J. S., M.A., University Observatory, Oxford. (1927.)
164. Hull, The Librarian, Botanical Department, University College. (1929.)
165. Humphrey, Dr C. J., United States Department of Agriculture, Soil Conservation Service, Stafford, Arizona, U.S.A. (1921.)
166. Hurst, Mr C. P., F.L.S., Landulph Rectory, Saltash, Cornwall. (1928.)
167. Ingold, Mr C. T., M.Sc., Ph.D., Department of Botany, University College, Leicester. (1935.)
168. Iowa, The Library, State University of Iowa, Library Annex, Iowa City, U.S.A. (1923.)
169. Iowa State College, Library, Ames, Iowa, U.S.A. (1927.)
170. Issatchenko, Professor B. L., Directeur du Jardin Botanique, Leningrad, Russia. (1923.)
171. John Crerar Library, 86 East Randolph Street, Chicago, Illinois, U.S.A. (1929.)
172. John Innes Horticultural Institution, Mostyn Road, Merton Park, London, S.W. 19. (1924.)
173. Johnson, Mr J. W. Haigh, M.Sc., F.I.C., F.L.S., Walton, nr. Wakefield. (1919.)
174. Jones, Mr G. H., M.A., Plant Protection Section, Ministry of Agriculture, Cairo, Egypt. (1922.)
175. Jørstad, Mr Ivar, Statsmykolog, Botanisk Museum, Oslo, Norway. (1923.)
176. Keay, Miss M. A., M.A., (Cape Town), Department of Agricultural Botany, The University, Reading. (1935.)
177. Keene, Miss K. N., B.Sc., West of Scotland Agricultural College, Blythwood Square, Glasgow. (1937.)
178. Keissler, Dr Karl, Direktor d. Botanischen Abteilung, Naturhistorisches Museum, Burgring 7, Wein 1/1, Austria. (1924.)
179. Kelly, Dr Howard A., 1418 Eutaw Place, Baltimore, Md., U.S.A. (1921.)
180. King, Miss M. E., B.A., The Botany School, Cambridge. (1935.)
181. Klika, Mr Bohumil, Hálkova, 37 Prague, Vrsovice 553 Czechoslovakia. (1926.)
182. Knight, Mr H. H., M.A., The Lodge, All Saints' Villas, Cheltenham. (1914.)

183. Kuala Lumpur, F.M.S., The Director of Agriculture, Straits Settlements, and Federated Malay States. (1930.)
184. Lamb, Mr I. M., British Museum (Natural History) London, S.W. 7. (1934.)
185. Lampitt, Mr L. H., D.Sc., F.I.C., Thornlea, Mount Park, Harrow, Middlesex. (1925.)
186. Leach, Mr R., B.A., Agricultural Department, Mlanje, Nyasaland. (1929.)
187. Leicester, The Museum, City of Leicester. (1923.)
188. Librarian, King's College, Newcastle-on-Tyne. (1928.)
189. Likhite, Dr Y. N., Department of Agriculture, Baroda State, India. (1936.)
190. Linder, Dr D., Farlow Herbarium, Harvard University, 20 Divinity Avenue, Cambridge, Mass., U.S.A. (1935.)
191. Line, Mr James, M.A., School of Agriculture, Cambridge. (1921.)
192. Linnean Society, The, Burlington House, Piccadilly, London, W. 1. (1919.)
193. Lloyd Library, The, 309 West Court Street, Cincinnati, Ohio, U.S.A. (1907.)
194. Loader, Miss F. M., B.Sc., Botanical Department, University College, Southampton. (1927.)
195. Lowndes, Mr A. G., M.A., F.L.S., Marlborough College, Marlborough, Wilts. (1922.)
196. Lütjeharms, Mr W. J., Department of Botany, University College of Orange Free State, Bloemfontein, S. Africa. (1930.)
197. McDonald, Mr J., D.F.C., B.Sc., F.L.S., Senior Plant Pathologist, P.O. Box 338 Nairobi, Kenya Colony, East Africa. (1923.)
198. McLennan, Dr Ethel I., Botanical Department, Melbourne University, Carlton, Victoria, Australia. (1926.)
199. Madras University Library, Senate House, Triplicane, Madras, South India. (1925.)
200. Maire, M. René, D.Sc., F.M.L.S., Professeur à la Faculté des Sciences de l'Université, Algiers, Algeria, N. Africa. (1907.)
201. Marsh, Mr R. W., M.A., Research Station, Long Ashton, Bristol. (1923.)
202. Masfield, Mr G. B., c/o Department of Agriculture, Entebbe, Uganda. (1932.)
203. Mason, Mr E. W., M.A., M.Sc., F.L.S., Imperial Mycological Institute, Ferry Lane, Kew, Surrey. (1921.)
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206. Mathias, Mr W. T., B.Sc., The University, Liverpool. (1938).
207. Mehta, Professor K. C., Ph.D., Department of Biology, Agra College, Agra, U.P., India. (1921.)
208. Melville, Mr R., B.Sc., Ph.D., 5 Courtway, Twickenham, Middlesex. (1933.)
209. Metcalfe, Mr C. R., B.A., Ph.D., Jodrell Laboratory, Royal Botanic Gardens, Kew, Surrey. (1926.)
210. Michigan Agricultural College Library, East Lansing, Michigan, U.S.A. (1924.)
211. Miller, Professor J. H., B.S., M.S., Ph.D., University of Georgia, Athens, Ga., U.S.A. (1930.)
212. Millidge, Mr P. H., 205 Carisbrooke Road, Newport, I.O.W. (1937.)
213. Missouri, The Botanical Garden, St Louis, Mo., U.S.A. (1902.)
214. Mitra, Mr M., M.Sc., Ph.D., D.I.C., Assistant Mycologist, Imperial Institute of Agricultural Research, Delhi, India. (1928.)
215. Miyabe, Dr Kingo, Professor Emeritus of Botany, Hokkaido Imperial University, Sapporo, Japan. (1919.)
216. Montague, Mrs A., Penton, Crediton, N. Devon. (1898.)
217. Montreal, Canada, Faculté des Sciences, Institut Botanique, Université de Montréal. (1932.)
218. Moore, Mr W. C., M.A., Ministry of Agriculture, Pathological Laboratory, Milton Road, Harpenden, Herts. (1922.)
219. Morgan, Dr G., Ashley-Hatton, Dyke Road Avenue, Brighton. (1928.)
220. Morris, Mr L. E., c/o Eton College, Windsor, Berks. (1924.)
221. Muller, Dr H. R. A., Institut voor Plantenziekten, Buitenzorg, Java. (1932.)
222. Murphy, Professor P. A., Sc.D., A.R.C.Sc.I., M.R.I.A., Department of Plant Pathology, Albert Agricultural College, Glasnevin, Dublin, N.W. 3. (1924.)
223. Murray, Mr G. H., F.E.S., Director of Agriculture, Rabaul, New Britain, Territory of New Guinea, via Australia. (1921.)
224. Muskett, Mr A. E., M.Sc., A.R.C.S., Queen's University, Belfast, Northern Ireland. (1923.)
225. Nannfeldt, Dr J. A., Sturegatan 11, Uppsala, Sweden. (1932.)
226. National Collection of Type Cultures, Curator, Lister Institute, Chelsea Gardens, London, S.W. 1. (1921.)
227. National Museum of Wales, Cardiff. (1924.)
228. Nash-Wortham, Mr J. R. H., Abingdon, Gatesden Road, Fetcham, Leatherhead, Surrey.

229. Nattrass, Mr R. M., B.Sc., (Agric.), Ph.D., Department of Agriculture, Nicosia, Cyprus. (1925.)
230. Nebraska, The Library, University of, Lincoln, Nebr., U.S.A. (1924.)
231. Nederlandsche Mycologische Vereeniging, c/o Dr A. J. P. Oort, Ericalaan 5, Wageningen, Holland. (1920.)
232. New York Botanical Garden, Bronx Park, New York, U.S.A. (1904.)
233. Nicholson, Mr W. E., F.L.S., 50 St Anne's Crescent, Lewes. (1913.)
234. Noel, Miss E. F., F.L.S., 37 Burnham Court, Queen's Road, London, W. 2. (1913.)
235. North Carolina, Library, University of, Chapel Hill, North Carolina, U.S.A. (1920.)
236. Nursery and Market Garden Industries' Development Society, Ltd., Experimental and Research Station, Cheshunt, Herts. (1922.)
237. O'Connor, Mr P., Ph.D., B.Sc., A.R.C.Sc.I., National Museum, Dublin. (1925.)
238. Ogilvie, Mr L., M.A., M.Sc., Research Station, Long Ashton, nr. Bristol. (1922.)
239. Oke, Mr Alfred William, B.A., F.G.S., F.L.S., 32 Denmark Road, Hove, Sussex. (1908.)
240. Ontario Agricultural College, Library, Guelph, Ontario, Canada. (1920.)
241. Osborn, Professor T. G. B., D.Sc., F.L.S., Botanical Department, The University, Oxford. (1910.)
242. Ottawa, Ontario, Canada, The Library, Geological Survey. (1926.)
243. Oyler, Miss E., Experimental and Research Station, Cheshunt, Herts. (1937.)
244. Padwick, Dr G. Watts, Imperial Agricultural Research Institute, New Delhi, India. (1936.)
245. Page, Miss W. M., M.Sc., Ph.D., 19 Ledam Buildings, Bourne Estate, Holborn, London, E.C. 1. (1921.)
246. Park, Mr M., Department of Agriculture, Peradeniya, Ceylon. (1929.)
247. Parke Davis and Co., Medical Research Library, P.O. Box 488, Detroit, Michigan, U.S.A. (1920.)
248. Parker, Professor C. S., Department of Botany, Howard University, Washington, D.C., U.S.A. (1932.)
249. Patrick, Miss S. H. M., Department of Botany, University, Oxford. (1937.) (P.A., Elmsleigh House, Stoughton Avenue, Leicester.)
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251. Peklo, Dr Jaroslav, Professor of Applied Botany, Bohemian Technical University, Charles Square, Prague II, Czechoslovakia. (1924.)
252. Pershouse, Mrs Stanley, Denhem Lodge, Yelverton, S. Devon. (1937.)
253. Perthshire Society of Natural Science, c/o J. F. Cumming, Esq., 12 Barossa Place, Perth. (1919.)
254. Petch, Mr T., B.A., B.Sc., North Wootton, King's Lynn, Norfolk. (1911.)
256. Pethybridge, Mr G. H., Ph.D., D.Sc., F.L.S., Seofen, Moreton End Lane, Harpenden. (1919.)
257. Peyronel, Dr Benjamino, R. Istituto Sup. Agrario e Forestale, Piazzale del Re, Firenze, Italy. (1932.)
258. Philadelphia, The Academy of Natural Sciences of Philadelphia, Nineteenth and The Parkway, Phil., U.S.A. (1925.)
259. Phillips, Dr H. H., 6 St John's Road, Penge, London, S.E. 10. (1923.)
260. Ping, Mr A. Wentworth, M.A., "St Olave's", Clifton, York. (1926.)
261. Pollard, Mr S. C., Jesus College, Oxford. (1937.)
262. Potter, Rev. M. C., Sc.D., M.A., F.L.S., Corley Croft, New Milton, Hants. (1896.)
263. Preston, Mr N. C., B.Sc., Harper Adams Agricultural College, Newport, Salop. (1920.)
264. Pretoria, South Africa, The Chief, Division of Plant Industry (91403), Department of Agriculture. (1922.)
265. Purdue University, Library, Agricultural Experiment Station, Lafayette, Ind., U.S.A. (1931.)
266. Ramsbottom, Mr J., O.B.E., Dr. Sc., M.A., Pres.L.S., British Museum, (Nat. Hist.), Cromwell Road, South Kensington, London, S.W. 7. (1910.)
267. Ray, Miss Anne, Penarwyn, Gorran Haven, Gorran, Cornwall. (1929.)
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269. Rea, Miss M. W., M.Sc., Salem House, Sydenham, Belfast, Northern Ireland. (1920.)
270. Rees, Mr John, M.Sc., Adviser in Agricultural Botany, University College, Cardiff. (1929.)
271. Reichert, Dr Israel, Jewish Agency for Palestine, Agricultural Experiment Station, P.O.B., 15 Rehoboth, Palestine. (1924.)
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273. Robinson, Mr E., 26 Burwood Avenue, Eastcote, Pinner, Middlesex. (1938.)

274. Rothamsted Experimental Station, Department of Mycology, Harpenden, Herts. (1923.)
275. Rutgers College and State University of New Jersey, Library, New Brunswick, New Jersey, U.S.A. (1922.)
276. St Paul, Minnesota, U.W.A., The Library, Department of Agriculture University Farm. (1920.)
277. Salisbury, Professor E. J., D.Sc., F.R.S., F.L.S., Botanical Department, University College, Gower Street, London, W.C. 1. (1921.)
278. Salmon, Professor E. S., F.L.S., South-Eastern Agricultural College, Wye, Kent. (1922.)
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283. Seth, Mr N. L., B.Sc., Ph.D., D.I.C., Agricultural College, Mandalay, Burma. (1930.)
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288. Smith, Professor Noel J. G., Ph.D., B.Sc., Botany Department, Rhodes University College, Grahamstown, S. Africa. (1924.)
289. Smith, Mr Rupert, 38 Greenhill Gardens, Edinburgh. (1927.)
290. South London Botanical Institute, 323 Norwood Road,ulse Hill, London, S.E. 24. (1921.)
291. Stakman, Professor E. C., University of Minnesota, Department of Agriculture, University Farm, St Paul, Minn., U.S.A. (1922.)
292. Statham, Miss E. M., 2 Westbrook Road, Blackheath, London, S.E. 3. (1926.)
293. Stationery Office, H.M., Superintendent of Publications, Book Dept., Westminster, S.W. 1. (4 subscriptions.) (1920.)
294. Stephens, Miss E. L., B.A., Department of Botany, University of Cape Town, South Africa. (1928.)

295. Stephens, Miss F. L., M.Sc., Department of Botany, British Museum (Natural History), Cromwell Road, South Kensington, London, S.W. 7. (1930.)
296. Steven, Mr W. F., Imperial Mycological Institute, Ferry Lane, Kew, Surrey. (1937.)
297. Steyaert, M. R. L., Ing. A.I.Gx., Laboratoire Bambesa, Uele, Belgian Congo. (1931.)
298. Stiles, Professor W., Sc.D., F.R.S., Department of Botany, The University, Edgbaston, Birmingham. (1936.)
299. Stirrup, Mr H. H., M.Sc., Midland Agricultural College, Sutton Bonnington, Loughborough. (1922.)
300. Storey, Mr J. H., M.A., Ph.D., East African Agricultural Research Institute, Amani, Tanganyika Territory, East Africa. (1922.)
301. Sutherland, Mr G. K., M.A., D.Sc., F.L.S., The Moorings, Rosemary Hill, Streetly, Sutton Coldfield, Birmingham. (1914.)
302. Swanton, Mr E. W., M.B.E., A.L.S., Educational Museum, Haslemere, Surrey. (1899.)
303. Swedish Academy of Sciences, Royal, Stockholm, Sweden. (1919.)
304. Sydney, Australia, The Librarian, University of. (1922.)
305. Sydow, Herr H., Luitpoldstrasse 33, Berlin, W. 30, Germany. (1931.)
306. Tabor, Mr R. J., B.Sc., F.L.S., Botanical Department, Imperial College of Science, South Kensington, London, S.W. 7. (1914.)
307. Tennessee, University of, Agricultural Experiment Station, Library, Knoxville, Tennessee, U.S.A. (1926.)
308. Tervet, Mr I. W., B.Sc., Department of Plant Pathology, University Farm, St. Paul, Minn. U.S.A. (1933.)
309. Tetley, Miss U., Quarry Garth, Windermere, Westmorland. (1929.)
310. Tomkins, Mr R. G., M.A., Ph.D., Trinity College, Cambridge. (1925.)
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312. Tunstall, Mr A. C., Tocklai Experimental Station, Cinnamara, P.O., Assam, British India. (1933.)
313. Vaheeduddin Syed, H. E. H. The Nizam's Government, Department of Agriculture, Hyderabad-Deccan, India. (1934.)
314. Vanterpool, Mr T. C., M.Sc., Botanical Department, University of Saskatchewan, Saskatoon, Canada. (1929.)
315. Venkatarayan, Mr S. V., Senior Assistant Mycologist, Agricultural Department, Bangalore, S. India. (1935.)

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317. Wakefield, Miss E. M., M.A., F.L.S., Herbarium, Royal Botanic Gardens, Kew. (1911.)
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319. Wales, University College of, Librarian, Botanical Department, Aberystwyth, North Wales. (1927.)
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326. Washington, Library, State College of, Pullman, Washington, U.S.A. (1924.)
327. Waterston, Mr J. M., B.Sc., Mycologist, Department of Agriculture, Paget East, Bermuda. (1934.)
328. Watson, Mr W., D.Sc., A.L.S., Cedene, Cheddou Road, Taunton. (1933.)
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333. Whetzel, Professor H. H., M.A., New York State College of Agriculture, Cornell University, Ithaca, N.Y., U.S.A. (1914.)
334. Whitaker, Mr F. Owen, 51 Grosvenor Avenue, Carshalton, Surrey. (1921.)
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342. Wiltshire, Mr S. P., D.Sc., Imperial Mycological Institute, Ferry Lane, Kew, Surrey. (1920.)
343. Wisconsin, The Library, University of, Madison, Wis., U.S.A. (1923.)
345. Wolf, Mr B. L., N.D.A., Cornwall Buildings, 45 Newhall Street, Birmingham. (1923.)
346. Wood, Mr F. C., The Rest, Franklin Road, Durrington, Worthing. (1935.)
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348. Woodward, Mr R. C., Ph.D., Imperial Chemical Industries, Ltd., Millbank, London, S.W. 1. (1924.)
349. Woolhope, The Naturalists' Field Club, Hereford. (1896.)
350. Worcestershire Naturalists' Field Club, Hereford. (1921.)
351. Wormald, Mr H., D.Sc., A.R.C.S., Research Station, East Malling, Kent. (1921.)
352. Wyatt-Smith, Mr J., Wadham College, Oxford. (1938.)
353. Yale University, Library, New Haven, Connecticut, U.S.A. (1930.)
354. Yeoman, Mr J. B., M.D., F.R.S.C., Norton, Wirral, Cheshire. (1934.)
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356. Zundel, Dr G. L. I., Botany Building, Pennsylvania State College, State College, Pa., U.S.A. (1929.)
357. Zürich, Switzerland, Botanical Garden and Museum, c/o Dr A. U. Däniker. (1921.)
358. Zürich, Institut für Spezielle Botanik der Eidg. Techn. Hochschule. (1928.)

*Members are requested to report any errors or omissions in the list to
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